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Alkylphenols Category SECTION TWO Ortho-substituted Mono-alkylphenols

Chemical Right-to-Know Initiative
HPV Challenge Program

Submitted to US EPA by: Schenectady International 2750 Balltown Road New York 12309

USA

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: not known.

RESULTS

Melting point: 14℃.

CONCLUSIONS

The test substance has a melting point 14°C.

DATA QUALITY

Not a GLP study.

Information taken from a literature search covering appropriate databases.

Purity of the test substance / decomposition: information not available.

REFERENCES

Ashford, R.D. Ashford's Dictionary of Industrial Chemicals. London, England: Wavelength Publications Ltd., 1994, p. 159.

2.1 BOILING POINT

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1993.

RESULTS

Boiling point: 224°C @ 760 mm Hg.

CONCLUSIONS

The test substance has a boiling point of 224° C.

DATA QUALITY

Not a GLP study.

Information taken from a technical datasheet.

Purity of the test substance given as typically 98.5 % minimum.

Decomposition: information not available.

REFERENCES

Schenectady International Inc. Technical Datasheet (11/93)

3.1 VAPOR PRESSURE

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 0.0173 mm Hg @ 25°C (2.31 Pa).

CONCLUSIONS

The test substance has a calculated vapor pressure of 0.0173 mm Hg @ 25℃.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 3.46 at 25°C.

CONCLUSIONS

The test substance has a calculated log Kow of 3.46 at 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

KOWWIN reported an experimental log Kow of 3.27 in its database (Hansch & Leo, 1985).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: calculation using a log Kow value of 3.46.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 319 mg/l @ 25°C

CONCLUSIONS

The water solubility of the test substance is 319 mg/l @ 25°C.

DATA QUALITY

Calculation.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: 44.1×10^{-12} cm³/molecule-sec.

Half-life: 2.91 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.91 hours.

DATA QUALITY

Calc ulation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:	150
Data temperature (°C):	25
Water solubility (mg/l):	319
Vapor pressure (Pa):	2.31
Log Kow:	3.27
Melting point (°C):	14

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	7.54
Soil	56.7
Water	34.4
suspended sediment	0.039
fish	0.0032
Sediment	1.26

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days/weeks

Ultimate biodegradation in weeks.

This material is may be readily biodegradable. This material is considered to be inherently biodegradable, and would aerobically biodegrade.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 2.78 mg/l

ChV (30 day) 0.41 mg/l

ChV (90 day) 0.040 mg/l

Remark: log Kow used 3.46 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 2.78 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 3.81 mg/l

ChV (96hr) 0.89 mg/l

Remark: log Kow used 3.46 (calculated value

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 3.81 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

Purity: Unknown

METHOD

Acute Toxicity to Non-Standard Species

Test type: unknown

GLP (Y/N): No data

Year study performed: 1981

Species: Crangon septemspinosa (Shrimp)

Analytical monitoring: No data

Exposure period: 96 hours

RESULTS

LC50 (96hr) 1.3 mg/l

The LC50 (96 hour) was determined to be 1.3 mg/l.

CONCLUSIONS

The LC50 (96 hour) was determined to be 1.3 mg/l.

DATA QUALITY

Unknown

REFERENCES

McLeese et al (1981) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 2.0 mg/l

ChV (21day) 0.30 mg/l

Remark: log Kow used 3.46 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 2.0 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

OECD Guidelines No. 401

GLP: (Y)

Year study performed: 1991

Species: Sprague-Dawley strain rat

Sex: Males and females

No of animals per sex per dose: Five males and five females

Vehicle: The test material was prepared as a solution in Arachis oil B.P.

Route of administration: Oral, gavage

Remarks: At the start of the study the animals were five to eight weeks old. A range-finding study was performed using pre-selected dose levels to determine the highest dose level that caused no deaths.

RESULTS

Range finding study:

hour of treatment. Increased salivation, ptosis and coma were noted before death.

Animals treated with test material in arachis oil B.P at dose levels of 5000 and 2000 mg/kg were found dead during the day of dosing or one day after treatment. Signs of systemic toxicity included lethargy, loss of righting reflex, decreased respiratory rate, ptosis, ataxia and hunched posture. Animals treated with 200 mg/kg bodyweight appeared normal throughout the study.

Based on this information, a dose level of 2000 mg/kg bodyweight in arachis oil was selected for the main study.

Main study:

There were no deaths. Lethargy was noted in all animals up to one hour after dosing. All animals in the main study appeared normal two hours after dosing and for the rest of the study and showed expected gain in body weight. No abnormalities were noted at necropsy.

CONCLUSIONS

The acute oral median dose (LD50) of the test material, in the Sprague-Dawley strain rat was found to be greater than 200 mg/kg body weight, but less than 2000 mg/kg body weight. The test material is classified as harmful and the symbol Xn and the risk phrase R22 "Harmful if swallowed" are therefore required according to the EEC labelling regulations.

DATA QUALITY

The study was conducted to OECD guidelines and GLP.

REFERENCES

Phenol 2-(1-methylpropyl): Range Finding Acute Oral Toxicity Test in the Rat –Test report by Sandoz Chemical td. Study Ref 47/1560. Issued by Safepharm Laboratories, P.O Box No. 45, Derby, DE1 2BT, U.K.

Study Ref: 47/1560

OTHER - ADDITION

An additional Rat Oral LD50 was 2700 mg/kg bw (NTP Chemical Repository)

NTP Chemical Repository as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

13.2 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

METHOD

OECD Guidelines No. 404

GLP: (Y)

Year study performed: 1991

Species: New Zealand White Rabbits

Sex: Males and females

No of animals per sex per dose: Two males and four females

Vehicle: 0.5 ml of the test material was introduced under a patch.

Remarks: A group of three rabbits were given a single 4-hour exposure semi-occluded application of the test material to the intact skin. The animals were observed for fourteen days after the day of the dosing. A further three animals were exposed to the test material for 3 minutes.

RESULTS

4-hour exposure period:

Severe erythema and edema was noted at all skin sites one hour after patch removal. Evaluation of the erythema and edema was not possible at the 24, 48, 72-hour and 7-day observations due to other adverse reactions. These included hemorrhage of the dermal capillaries, hardened dark brown/black colored scabs, blanching, well-defined of moderate erythema surrounding the treatment site, undulating scabs, scabs lifting at edges to reveal either dried blood or light brown-colored scabs with small areas of dried blood. These reactions were considered to be indicative of dermal corrosion.

3-minute exposure period:

Well-defined or moderate to severe erythema was noted at all treated skin sites one, 24, 48 and 72-hours after patch removal. Moderate to severe edema was noted at all treated skin sites one hour after patch removal. Slight to severe edema was noted at the 24-hour observation with very slight to moderate edema at the 72-hour observation. Very slight to mild edema was noted at two treated skin sites at the 7-day observation. Evaluation of the erythema and edema was not possible at one treatment site at the 72-hour observation and all treated sites at the 7-day observations due to other adverse reactions. These included hemorrhage of the dermal capillaries, light brown discoloration of the epidermis, loss of skin elasticity, thickening of the skin, hardened dark brown/black-colored scab, well-defined erythema surrounding the treated site, hardened light brown-colored scab, reduced regrowth of fur and desquamation.

CONCLUSION

The test material was classified as corrosive according to the EEC labelling regulations. The symbol C and risk phrase R34 "Causes burns" are required.

DATA QUALITY

Study conducted to OECD guidelines and GLP

REFERENCES

Phenol 2-(1-methylpropyl): Acute Dermal Irritation Test in the Rabbit, Test Report by Sandoz Chemicals. Study Ref: 47/1561. Issued by Safepharm Laboratories, P.O. Box No. 45, Derby, DE1 2BT, U.K.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

o-sec-Butylphenol

CAS No. 89-72-5

The substance used for this report was 98%.

METHOD

Guideline followed: Ames test Zeiger and Drake 1980; Haworth et al, [1983]

Type: The Salmonella preincubation assay System of testing: Bacterial

GLP: No data available, but given the date, author and sponsoring agency GLP is likely.

Year study performed: 1985

Species/strain: Salmonella typhimurium, TA 1535, TA 1537, TA 98 and TA 100.

Metabolic activation: Aroclor 1254 induced rat or hamster liver 10% S-9 fraction

Concentrations tested: 0, 3, 10, 16, 33, 100 and 166 µg/plate

Remarks: Chemicals were tested and evaluated as coded samples (aliquots) using *Salmonella* strains treated in the presence and absence of metabolic activation with the test material dissolved in DMSO. At least five dose levels of the chemicals were tested, with three plates per dose level. All assays were repeated. Concurrent solvent and positive controls were tested with and without the metabolic activation systems.

The following mutagens were used as concurrent positive controls: sodium azide for TA 1535 and TA 100, 4-nitro-o-phenylenediamine for TA 98 and 9-aminoacridine for TA 97 and TA 1537; 2-aminoanthracene was used with all strains with hamster and rat liver metabolic activation systems. The dose levels used by all three laboratories have been reported elsewhere [Haworth et al, 1983]. Three of the mutagens, 9-aminoacridine hydrochloride H₂O, 4-nitro-o-phenylenediamine and tris (1,3-dichcloro-2-propyl)phosphate were positive controls which were sent coded to each laboratory. Potassium chloride was a coded negative control.

The criteria used for data evaluation were based on a) mutagenic a dose related increase in number of revertants over background even if the increase was less than twofold, b) non mutagenic response when no increase in the number of revertants was elicited by the chemical c) equivocal when there was an absence of clear-cut-dose-related increase in revertants.

RESULTS

In this study o-sec butylphenol did not induce a dose-related increase in mutant colonies over DMSO in any strain in the absence or presence of metabolic activation. Slight clearing of background lawn was observed at 166 μ g/plate in strains TA 100, TA 1537 and TA 98 without metabolic activation.

CONCLUSIONS

The test substance is not genetically active in the Ames Salmonella typhimurium assay.

Lab: SRI

DATA QUALITY

No mention of GLP, but given the date, author and sponsoring agency GLP is likely.

REFERENCES

Salmonella Mutagenicity Tests: II. Results from the Testing of 270 Chemicals

OTHER

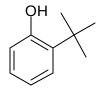
Study Ref: Environmental Mutagenesis Volume 8, Supplement 7:1-119 (1986)

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2-tert-Butylphenol



CAS No. 88-18-6

METHOD

Method/guideline followed: information not available.

GLP (Y/N): No.

Year study performed: Not known.

RESULTS

Melting point: -6.8°C.

CONCLUSIONS

The test substance has a melting point of -6.8°C.

DATA QUALITY

Not a GLP study.

Information taken from standard reference book (peer reviewed).

Purity of the test substance / decomposition: information not available in reference.

REFERENCES

Lide, D.R. (ed.). CRC Handbook of Chemistry and Physics. 75th ed. Boca Raton, F1: CRC Press Inc., 1994-1995., p. 3-255.

2.1 BOILING POINT

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: information not available in reference.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Boiling point: 223°C.

Pressure: 760 mm Hg.

CONCLUSIONS

The test substance has a boiling point of 223° C.

DATA QUALITY

Not a GLP study.

Information taken from standard reference book (peer reviewed).

Purity of the test substance / decomposition: information not available in reference.

REFERENCES

Lide, D.R. (ed.). CRC Handbook of Chemistry and Physics. 75^{th} ed. Boca Raton, F1: CRC Press Inc., 1994-1995., p. 3-255.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): not known.

Year study performed: not known.

RESULTS

Vapor pressure: 0.09 mm Hg @ 25°C (12.0 Pa).

CONCLUSIONS

The test substance has a vapor pressure of : 0.09 mm Hg @ 25°C (12.0 Pa).

DATA QUALITY

Extrapolated value.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

SRC PhysProps database using data from Perry, R.H. and Green, D. (1984); Perry's Chemical Engineers' Handbook., 6^{th} ed. McGraw-Hill, New York.

OTHER

A vapor pressure of 14.85 Pa is quoted for 2-*tert*-butylphenol in Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: not known.

GLP (Y/N): not known.

Year study performed: 1995.

RESULTS

Log Kow: 3.31.

CONCLUSIONS

The test substance has a log Kow of 3.31.

DATA QUALITY

Information taken from Hazardous Substances Database (HSDB).

REFERENCES

Hansch, C., Leo, A., Hoekman. Exploring QSAR – Hydrophonic, Electronic and Steric Constants. Washington DC: American Chemical Society, 1995 as cited in HSDB Database.

OTHER

A pKa value of 10.28 is quoted in Schueuerman G; Sci Total Environ 109/110: 221-35 (1991). Original reference: Schultz, T.W. and Cajina-Quezada, M.; Structure-Activity Relationships for Mono Alkylated or Halogenated Phenols, Toxicol. Lett., 37 (1987) 121-130

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: calculation using a calculated log Kow value of 3.42.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 343.9 mg/l @ 25℃.

CONCLUSIONS

The test substance has a calculated water solubility of 343.9 mg/l.

DATA QUALITY

Calculation.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v1.33.

OTHER

A pKa value of 10.28 is quoted in Schueuerman G; Sci Total Environ 109/110: 221-35 (1991). Original reference: Schultz, T.W. and Cajina-Quezada, M.; Structure-Activity Relationships for Mono Alkylated or Halogenated Phenols, Toxicol. Lett., 37 (1987) 121-130

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: 40.6×10^{-12} cm³/molecule-sec.

Half-life: 3.16 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 3.16 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

COMMENT

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:150Data temperature (°C):25Water solubility (mg/l):344Vapor pressure (Pa):12Log Kow:3.31Melting point (C):-6.8

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	27.0
Soil	46.3
Water	25.6
suspended sediment	0.032
fish	0.0026
Sediment	1.03

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days - weeks

Ultimate biodegradation in weeks - months.

This material may be readily biodegradable. This material is considered to be inherently biodegradable, and would aerobically biodegrade.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

Purity of the test substance: mixture containing 2.273% of 2-tert-butylphenol.

METHOD

Method: OECD Guideline No. 203
US EPA 40 CFR Part 797.1400
Test type: Static

GLP: (Y)

Year study performed: 1992

Species/strain: Fathead minnow Pimephales promelas

Test substance preparation: Water used for acclimation of test organisms and for all toxicity testing was dechlorinated tap water. The test substance was added directly to dilution water to formulate test media without the use of a solvent.

Analytical monitoring: None, nominal concentrations were used.

Exposure period: 96 hours

Statistical methods: Results of the toxicity test were interpreted using standard statistical techniques (the probit method). Computer methods (Stephan, 1983) were used to calculate LC50 values.

Deviations from protocol:

The hardness of the dilution water was greater than 50 mg/l and fish were not acclimated to the water hardness employed during the definitive test. These protocol deviations did not, in the author's opinion, affect the outcome of the toxicity test. No other deviations were made from the protocol.

Test conditions: The test was conducted as an acute screening test. Ten fathead minnows were indiscriminately distributed to a single replicate of each concentration. A 16 hour light and 6 hour dark photoperiod was maintained. Aeration was not required to maintain dissolved oxygen concentration above acceptable levels.

Control:

Temperature $21.1-22.1^{\circ}\text{C}$ Dissolved oxygen 75 -102% saturation pH 7.8 -8.2Conductivity 600-660 µohms/cm Hardness 176 mg/l CaCO₃

Test treatment:

Temperature $21.5-22.3^{\circ}\text{C}$ Dissolved oxygen $51-102^{\circ}\text{m}$ saturation pH 7.8-8.3Conductivity 600-670 µohms/cm Hardness 176 mg/l CaCO₃

Remarks: The first acute screening test was conducted for 96 hours. Nominal concentrations were 0, 1, 10, 100 and 1,000 mg/l. The concentrations were formulated by the addition of the test substance directly to the dilution water without the use of a solvent. The test was repeated because dilution water had an incorrect hardness.

RESULTS

Nominal concentrations: 0, 1, 10, 100 and 1,000 mg/l

Remarks:

During the test all non-control test vessels contained insoluble material that was stuck to the plastic weight boat (weight boats were placed in the test vessels) and on the surface in all test vessels containing the test substance.

LC50 (96hr): 680 mg/l

Nom. concn.	Percentage mortality				Percentage affected					
(mg/l)	0	24	48	72	96	0	24	48	72	96
Control	0	0	0	0	0	0	0	0	0	0
1	0	0	0	0	0	0	0	0	0	0
10	0	0	0	0	0	0	0	0	0	0
100	0	0	0	0	10	0	0	0	0	0
1000	0	0	50	50	60	0	100	50	50	40

All affected fish exposed to 1,000 mg/l were lethargic, gasping, exhibited erratic swimming, and/or were dark in color at 24 to 96 hours.

CONCLUSIONS

Exposure of test organisms to the test substance, a mixture containing 2.273% of 2-*tert*-butylphenol, resulted in a 96 hour LC50 of 680 mg/l (equivalent to 15.5 mg 2-*tert*-butylphenol/l).

DATA QUALITY

Method follows OECD guideline

GLP study

REFERENCES

Acute Toxicity of a Mixture of 2-tert-Butylphenol to the Fathead Minnow (*Pimephales promelas*) Results of Range finding test. Document ID 8690000471s TR Wilbury Labs Inc . Massachusetts 01945

10.2 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Fish

Exposure period: 96 hours, 30 days, 90 days

RESULTS

LC50 (96hr) 2.935 mg/l

ChV (30 day) 0.433 mg/l

ChV (90 day) 0.041 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 2.935 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Estimation performed for this HPV submission

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 4.111 mg/l

ChV (96hr) 0.935 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be $4.111\ mg/l$.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Estimation performed for this HPV submission

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6



Purity: Unknown

METHOD

Acute Toxicity to Non-Standard Species

Test type: unknown

GLP (Y/N): No data

Year study performed: 1981

Species: Crangon septemspinosa (Shrimp)

Analytical monitoring: No data

Exposure period: 96 hours

RESULTS

LC50 (96hr) 2.4 mg/l

The LC50 (96 hour) was determined to be 2.4 mg/l.

CONCLUSIONS

The LC50 (96 hour) was determined to be 2.4 mg/l.

DATA QUALITY

Unknown

REFERENCES

McLeese et al (1981) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 2.118 mg/l

ChV (21day) 0.318 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 2.118 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Estimation performed for this HPV submission

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

Purity of test substance: 99.96% by GLC

METHOD

Method: OECD Guideline No. 401

GLP: Data not available

Year study performed: 1990

Species/strain: Fischer 344 rats

No. of animals per sex per dose: 5 males and 5 females

Vehicle: Corn oil

Route of administration: Oral, gavage

Dose level: 474, 664, 930, 2551 and 5000 mg/kg

Remarks: A preliminary test utilising groups of one male and one female rat treated at 100, 500 and 1500 mg/kg indicated that the acute median lethal oral dose (LD50) was between 500 and 1500 mg/kg.

RESULTS

LD50 789 mg/kg (95% confidence interval 653 to 1017), slope 8.9, males and females LD50 789 mg/kg (95% confidence interval 695 to 914), slope 8.9, males and females combined.

Number of deaths at each dose level:

Dose (mg/kg)	Cumulative mortality	
	Male	Female
474	0/5	0/5
664	1/5	1/5
930	4/5	4/5
2551	5/5	5/5
5000	5/5	5/5

There were deaths among rats dosed at 664 mg/kg and above and no rat survived oral administration of the test substance at 2551 and 5000 mg/kg. The majority of deaths occurred during Day 1 (5000 mg/kg) or Day 2 (930 and 2551 mg/kg) but single animals dosed at 664 and 930 mg/kg were found dead on Day 3 and a single male was killed on humane grounds on Day 10.

Clinical signs: Lachrymation, abasia/ataxia, and prostration. The time of onset of these clinical signs was inversely related to the administered dose. A hunched posture and lethargy were common at the lower and intermediate dose levels, particularly at 664 mg/kg. An unkempt appearance and/or yellow staining of the anogenital zone developed in the majority of rats surviving to Day 2. Among the animals dosed at 5000 mg/kg there were incidences of pallor or a darkened appearance of the eye, cyanosis and wheezing. Coma was observed at dose levels of 664 mg/kg and was always followed by death. There were isolated cases of diarrhoea, tachypnoea, hypothermia, tremor, salivation, piloerection, epistaxis, periorbital encrustation and swelling or opacity of the eye.

Most clinical signs were first apparent within 4 hours of dosing. Recovery of rats surviving treatment, as judges by external appearance and behaviour was advanced by Day 3 and completed by Day 11.

All surviving rats had gained weight relative to their Day 1 body weights by the end of the 14 day observation period.

Necropsy findings:

An unkempt appearance, corneal opacity and either pallor or a darkened appearance of the eyes were commonly noted. The principal internal macroscopic abnormalities revealed exaggerated hepatic lobular pattern, darkened liver, darkened spleen, renal pallor and/or a granular appearance of the kidneys and inflammation with abnormal content of the gastrointestinal tract. No significant lesions were found among the rats terminated on Day 15.

CONCLUSIONS

Acute oral LD50 of 2-tert-butylphenol in fasted rats of both sexes was found to be 789 mg/kg.

DATA QUALITY

Method follows OECD Guideline

REFERENCES

O-tert-butylphenol – Acute Oral Toxicity.
Document ID 86960000557
Shell Research Ltd, Sittingbourne Research Centre, Sittingbourne, Kent, England

13.2 ACUTE TOXICITY DERMAL

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

Purity of the test substance: 99.96% by GLC

METHOD

Method: OECD Guideline No. 402

GLP: Data not available

Year study performed: 1990

Species/strain: Fischer 344 rats

No of animals per sex per dose: 5 males and 5 females

Vehicle: Substance administered as such

Route of administration: Dermal

Dose levels: 1020, 1420 and 2000 mg/kg (males) and 520, 720, 1020 and 2000 mg/kg

(females)

Remarks: A single dose of the undiluted test material was applied to the rat skin and held in place with a lint dressing covered with waterproof adhesive tape. Following a 24-hour exposure the dressings were removed, the skin washed with warm dilute detergent solution and dried.

RESULTS

LD50 1373 mg/kg (95% confidence interval 1124 - 1473), slope 40.9, males LD50 705 mg/kg (95% confidence interval 574 - 756), slope 41.0, females

There were deaths on Day 2 to 5.

Clinical signs:

Hematuria was apparent in all rats except three females that were among the first decedents. There were no other clinical signs among rats dosed at 520 mg/kg.

The reactions to treatment at higher dose levels were lethargy and prior to death lachrymation, pale eyes, hypothermia, prostration and coma. Isolated cases of skin pallor, periorbital encrustation, hunched posture, unkempt appearance and yellow staining of the anogenital zone were also observed. Recovery from these clinical signs was advanced by Day 2 and, with one exception, was complete by Day 7.

Losses of body weight or minimal body weight gains were recorded for all rats surviving at completion of the first week of the observation period. All surviving rats had gained weight relative to their Day 1 body weights at the end of the 14 day observation period.

After removal of the occlusive dressing on Day 2, the treated skin showed inflammation, a brown wrinkled appearance or a chemical burn. Scab formation followed between Days 6 and 10 and this persisted at termination on Day 15.

Necropsy findings:

Necropsy findings of decedents included darkened appearance and potechiation of thymus, soft brain, lung congestion, pallor and exaggerated lobular pattern of the liver, dark spleen, pallor of the renal cortex or darkening of the renal medulla, inflammation and abnormal content of the urinary bladder, inflammation of stomach and abnormal gastrointestinal contents. Four rats killed at Day 14 showed exaggerated hepatic lobular pattern.

Findings of inflammation, discoloration, scab formation and subcutaneous congestion or inflammation at the dermal test sites were consistent with in-life observations.

CONCLUSION

This material has a male rat LD50 of 1373 mg/kg.

This material has a female rat LD50 of 705 mg/kg.

As there was a clear sex-related difference in the toxicity of 2-tert-butylphenol, no LD50 value for rats of both sexes were computed.

DATA QUALITY

Method follows OECD Guideline.

REFERENCES

O-tert-butylphenol – Acute Dermal Toxicity
Document ID 86960000557
Shell Research Ltd, Sittingbourne Research Centre, Sittingbourne, Kent, England

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY)

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

Purity of the test substance: Data not available however, identity was confirmed by comparing an infra-red spectrum of the test substance with a published spectrum of 2-tert-butylphenol.

METHOD

Method: Ames test

Type: Reverse mutation assay

System of testing: Bacterial, Plate Incorporation Assay was used.

GLP: (Y)

Year study performed: 1989

Species/strain: Salmonella typhimurium strains TA98, TA100, TA1535, TA1537 and TA1538.

Ecsherichia coli WP₂ uvrA pKM10

Metabolic activation: Aroclor 1254 induced rat liver 10% S-9 fraction.

Concentrations tested: 31.25, 62.5, 125, 250, 500, 1000, 2000 and 5000 µg/plate.

Test design: The solutions of the test substance in acetone were added to top agar mix and were tested both in the presence and absence of S-9 mix. All tests were carried out in triplicate. The replicate assays were carried out on different days in order to confirm the reproducibility of the results. All cultures were incubated at 37°C for 48-72 hours before the revertant colonies were counted.

Remarks: The test compound formed an oily film on the surface of the top agar at 500 μ g/plate and above, showing that it was not miscible in the aqueous test system at these concentrations.

The addition of 2500 μ g/plate per ml of the test substance (equivalent to approx. 5000 μ g/plate) caused the pH of the medium to change from 7.31 to 7.42.

The following mutagens were used as concurrent positive control:

Bacterial strain	Positive control	
	(without S-9)	(with S-9)
E. coli WP 2 uvrA pKM10	Potassium dichromate	Benzo (a) pyrene
S. typhimurium <i>TA1535</i>	Sodium azide	2-Aminoanthracene
S. typhimurium TA1537	9-Aminoacridine	Neutral red
S. typhimurium <i>TA1538</i>	2-Nitrofluorene	Benzo (a) pyrene
S. typhimurium <i>TA98</i>	2-Nitrofluorene	Benzo (a) pyrene
S. typhimurium TA100	Sodium azide	Benzo (a) pyrene

RESULTS

In this study, the test substance did not increase the reverse mutation frequency in any of the bacterial strains, in either presence or absence of metabolic activation. Microscopic evaluation of the background lawn showed evidence of cytotoxicity in all bacterial tester strains both in the presence and absence of S-9 mix.

CONCLUSIONS

2-tert-butylphenol is not genotoxic in the selected bacterial tester strains.

DATA QUALITY

GPL study

REFERENCES

Bacterial Mutagenicity Studies with *Ortho-Tert*-Butylphenol Document ID 8690000558
Shell Research Ltd. Sittingbourne Research Centre, Sittingbourne, Kent, England

OTHER TUMOUR INHIBITORY EFFECTS OF PHENOLIC COMPOUNDS ON BENZO(A) PYRENE-INDUCED NEOPLASIA

TEST SUBSTANCE

2-tert-Butylphenol

CAS No. 88-18-6

Purity of the test substance: 99%

METHOD

Method: See below

GLP: Data not available

Year study performed: 1980

Species/strain: Mouse ICR/Ha strain

Route of administration: Oral, dietary

Exposure period: 38 days

Dose/concentration: 0.03 mmol/g (= 4,500 ppm in diet)

Sex: Females

No of animals per dose: 20 in each of two experiments.

Frequency of treatment: continuous in the diet for 5.5 weeks. 1 mg benzo(a)pyrene (BP) by gavage, twice/week for 4 weeks, from 2^{nd} week of test diet administration.`

Positive control: BP as before, but no dietary addition of the test substance

Post exposure observation period: 16 weeks

Statistical methods: Student's T test was used to determine the statistical differences in the number of tumors per group between the control and treated groups, and the χ^2 test was used for the differences in percentage of tumour bearing animals in these groups.

Test conditions: The mice were placed on a diet containing the test substance or diet without additions (control group) when they were 9 weeks of age. On the eighth day, the mice were given the first of 8 doses (2 times a week for 4 weeks) of 1 mg of BP in 0.2 ml of corn oil. The experimental diets were

fed during the entire period of carcinogen administration and were discontinued 3 days after the last dose of carcinogen, at which time the mice were 98 days old. The mice were then fed pellets of rat chow, until they were killed at 211 days old and autopsied. Tumors of the forestomach were counted under a dissecting microscope. Tumors that were 1 mm or larger were recorded and checked histologically.

RESULTS

In the control group (animals given BP but no dietary addition of phenol), 90 - 100% of animals had neoplasms of the forestomach and the average number of tumours per animal was between 4.1 and 5.8. In the animals given dietary addition of 2-tert-butylphenol, 85 - 95% of animals had neoplasms of the forestomach, but the number of neoplasms per animal was reduced to about 60% of the controls, which was statistically significant.

CONCLUSION

Significant inhibitory activity of 2-tert-butylphenol was demonstrated in this mouse model, but the relevance of this to possible protection effects in humans is not known.

DATA QUALITY

Reliable, without restrictions

REFERENCES

Department of Laboratory Medicine and Pathology, University of Minnesota, Minneapolis. Cancer Research 40, 2820-2823. August 1980.

Alkylphenols Category SECTION THREE Para-substituted Mono-alkylphenols

Chemical Right-to-Know Initiative
HPV Challenge Program

Submitted to US EPA by:

Schenectady International

2750 Balltown Road

New York 12309

USA

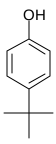
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No

Year study performed: 1992

RESULTS

Melting point: ca. 100°C.

CONCLUSIONS

The test substance has a melting point of ca. 100°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

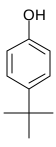
Sicherheitsdatenblatt Huels AG, 28.04.92 as cited in IUCLID database.

2.1 BOILING POINT

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: 1992.

RESULTS

Boiling point: 237°C at 101325 Pa (760 mm Hg).

CONCLUSIONS

The test substance has a boiling point of 237° C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

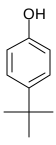
Sicherheitsdatenblatt Huels AG, 28.04.92 as cited in IUCLID database.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: 1994.

RESULTS

Vapor pressure: 0.5 Pa at 20°C.

CONCLUSIONS

The test substance has a vapor pressure of 0.5 Pa at 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

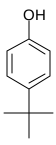
Huels AG: Produktinformation "p-tert-Butylphenol", Art.-Nr.: 001786; Ausgabe 01.08.1994 as cited in IUCLID database.

4.1.PARTITION COEFFICIENT

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: flask-shake method.

GLP (Y/N): No.

Year study performed: not known.

RESULTS

Log Kow: 3.31.

CONCLUSIONS

The test substance has a log Kow of 3.31.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance: information not available in database.

The flask-shake method is an acceptable technique and is described in EU, OECD and OPPTS test guidelines.

REFERENCES

Church, C., Hansch, C.: unpublished results cited in: Leo, A. et al. (1971): Chem. Rev. 71, 531, 537 – 538, 551, 555, 593. Cited in IUCLID database.

OTHER

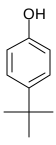
p-tert-Butylphenol has a pKa value of 10.39 (Serjeant, E.P., Dempsey, B. (1979) (Ionization Constants of Organic Acids in aqueous Solution, Pergamon, N.Y. as cited in the IUCLID database.)

5.1. WATER SOLUBILITY

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: 1990.

RESULTS

Water solubility: 0.8 g/l at 20 °C.

CONCLUSIONS

The water solubility of the test substance is 0.8 g/l at $20 \,^{\circ}\text{C}$.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance: information not available in database.

REFERENCES

Boeddeker, K.W. et al. (1990): J. Membr. Sci. 53, 143 – 158 as cited in IUCLID database.

OTHER

p-tert-Butylphenol has a pKa value of 10.39 (Serjeant, E.P., Dempsey, B. (1979) (Ionisation Constants of Organic Acids in aqueous Solution, Pergamon, N.Y. as cited in the IUCLID database.)

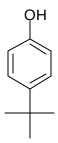
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): No.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $40.6 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 3.16 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 3.16 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

Direct aqueous photolysis: *p*-cresol in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

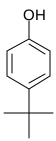
7.1 STABILITY IN WATER - ADDITION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4

Purity: 99%



METHOD

Test type: OECD TG 111

GLP (Y/N): Yes.

Year study performed: 1997

RESULTS

Stable at pH 4, 7, and 9 at 25 C.

CONCLUSIONS

Stable at pH 4,7, and 9 at 25 C.

DATA QUALITY

GLP Compliant

REFERENCES

MITI, Japan (1997) as cited in the SIDS Dossier on the HPV Phase-4 Chemical, p-tert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

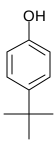
The software prediction program HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: calculation made for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:150Data temperature (°C):25Water solubility (mg/l):800Vapor pressure (Pa):0.50Log Kow:3.31Melting point (C):100

Environmental conditions: defaults used

RESULTS

Environmental compartment	Percentage of test substance	
Air	0.66	
Soil	63.0	
Water	34.9	
suspended sediment	0.044	
Fish	0.0036	
Sediment	1.40	

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

OTHER - ADDITION

Mackey Level III performed for the SIDS Dossier:

Media: air-biota-sediment-soil-water

Results:

COMPARTMENT	RELEASE – 100% to air	RELEASE – 100% to water	RELEASE – 100% to soil
Air	39.7 %	0.2 %	0.0 %
Water	23.3 %	95.3 %	0.4 %
Soil	35.9 %	0.2 %	99.6 %
Sediment	1.1 %	4.4 %	0.0 %

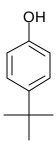
Section 3.3.2 in the SIDS Dossier on the HPV Phase-4 Chemical, p-tert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

9.1 BIODEGRADATION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Method/guideline followed: Directive EEC/92/69, Part II, C. 4A DOC Die Away Test

Test type: aerobic

GLP (Y/N): Yes.

Year study performed: 1992

Contact time: 28 days.

Inoculum: non-adapted, domestic, activated sludge.

Concentration: 10 mg/l (related to DOC).

RESULTS

Percentage degradation: 98% after 28 days

CONCLUSIONS

The test substance was readily biodegradable under the conditions of the test.

DATA QUALITY

GLP study.

The test concentration was well within the water solubility of p-tert-butylphenol.

REFERENCES

Huels report No. DDA-59, 1994 (unpublished) as cited in IUCLID database.

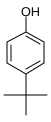
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

*p- tert-*butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance given as 99%.

METHOD

Acute Toxicity Test, based on US EPA procedures

Test type: Flow through toxicity test

GLP (Y/N): Data not available

Year study performed: 1975

Species: Freshwater fish - Pimephales promelas

Fish used for the test were 31 to 35 days old.

Analytical monitoring: Yes, no details available on IUCLID.

Exposure period: 96 hours

Test conditions:

Temperature: 24.6 ± 1.4 °C

pH 6.9 - 7.7

Dissolved oxygen: $4.6 - 8.8 \text{ mg O}_2/1$

Hardness (CaCO₃): 42.2 – 46.6 mg/l

RESULTS

LC50 (24hr) 6.21 mg/l

LC50 (48hr) 5.69 mg/l

LC50 (72hr) 5.26 mg/l

LC50 (96hr) 5.14 mg/l

Remarks: 5.44 mg/l caused some cases of spinal deformities; at 3.10 mg/l some fish were narcotised (no reaction to tapping the outside of the tank).

CONCLUSIONS

The LC50 (96hr) of the test substance is 5.14 mg/l.

DATA QUALITY

Information taken from IUCLID database.

GLP study: data not available.

REFERENCES

IUCLID Data Sheet

4.1 Acute/Prolonged Toxicity to Fish

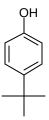
Study Ref.: Huels AG Marl

10.2 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

*p- tert-*butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance given as 99.5%.

METHOD

Acute Toxicity Test, OECD TG 203

Test type: Semi-static, open system toxicity test

GLP (Y/N): No.

Year study performed: 1995

Species: Oryzias latipes (Medaka)

Analytical monitoring: Yes, no details available.

Exposure period: 96 hours

Group of ten Medaka were exposed to nominal concentrations of 2.0, 3.0, 4.5, 6.8, and 10 mg/l. 100 mg/l DMSO and HCO-40 (4:1 weight ratio) was used as a solubiliser. 100 mg/l solubiliser and dechlorinated tap water were used as a control.

RESULTS

The LC50 (96 hour) was determined to be to be 5.1 mg/l with 95% confidence limits of 4.7 mg/l to 5.8 mg/l.

LC50 (24 hr): 5.1 mg/l

LC50 (48 hr): 5.1 mg/l

LC50 (72 hr): 5.1 mg/l

LC50 (96 hr): 5.1 mg/l

CONCLUSIONS

The LC50 (96hr) of the test substance is 5.1 mg/l.

DATA QUALITY

Information taken from SIDS Dossier.

REFERENCES

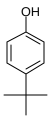
Environment Agency of Japan (1995) as cited in the SIDS Dossier on the HPV Phase-4 Chemical, ptert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

10.3 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance given as > 99%.

METHOD

Acute Toxicity Test

Test type: Semi-static, open system toxicity test

GLP (Y/N): Unknown

Year study performed: 1984

Species: *Pimephales promelas* (Fathead Minnow)

Analytical monitoring: Yes, no details available.

Exposure period: 96 hours

Fifty fathead minnows (25 per duplicate tank) were exposed to five test concentrations in flow-through aquaria (7.3 l at flow rate of 83 ml/min) for 96 hours.

Water for testing was obtained from Lake Superior. Age of fathead minnows was 31-35 days old. Water temperature of 24.5 C; dissolved oxygen of 7.4 mg/l; hardness of 44.9 mg/l as CaCO₃

Toxicity Tests were conducted together with 24 chemicals.

RESULTS

The LC50 (96 hour) was determined to be to be 5.1 mg/l.

LC50 (24 hr): 6.2 mg/l

LC50 (48 hr): 5.7 mg/l

LC50 (72 hr): 5.3 mg/l

LC50 (96 hr): 5.1 mg/l

CONCLUSIONS

The LC50 (96hr) of the test substance is 5.1 mg/l.

DATAQUALITY

Information taken from SIDS Dossier.

REFERENCES

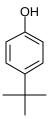
Holcombe G.W., Phipps, G.L., Knuth, M.L., and Felhaber, T. (1984) Environ. Pollut. (Ser. A) 35, 367-381, as cited in the SIDS Dossier on the HPV Phase-4 Chemical, p-tert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

10.4 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP (Y/N): No.

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 2.9 mg/l

ChV(30 day) 0.43 mg/l

ChV (90 day) 0.041 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 2.9 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

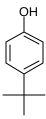
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE) - ADDITION

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: 99.5%

METHOD

Algal Biomass Test OECD TG 201 (1984)

Test type: Static, open-system

GLP (Y/N): No.

Year study performed: 1995

Species: Selenastrum capricornutum ATCC22662

Analytical monitoring: Yes, but no data

Exposure period: 72 hours

The EC50 value for biomass change (% inhibition) was calculated based on five nominal concentrations (9.53, 17.2, 30.9, 55.6, and 100 mg/l). Minimal amount of Tween 80 – acetone (1:1) or DMSO-HCO 40 (9:1) is used as a stabiliser.

RESULTS

Based on biomass:

EC50 (72 hr) 22.7 mg/l

NOEC 9.53 mg/l

LOEC 17.2 mg/l

CONCLUSIONS

The EC50 (72hr) of the test substance was found to be $22.7\ mg/l$.

DATA QUALITY

Included in SIDS Dossier

REFERENCES

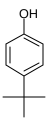
Environment Agency of Japan (1995) as cited in the SIDS Dossier on the HPV Phase-4 Chemical, ptert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: information not available. Re-crystallised before testing.

METHOD

Algal Growth Inhibition Test	
Test type: Data not available	
GLP (Y/N): Data not available	
Year study performed: 1971	

Species: Chlorella vulgaris

Analytical monitoring: No data

Exposure period: 6 hours

Test conditions:

Temperature: 36.5°C

RESULTS

EC50 (6hr) 22.2 - 34.4 mg/l

Remarks: The EC50 value depended on the method of biomass determination:

photometric extinction at 680 nm: EC50 = 22.2 mg/l

photometric extinction at 750 nm: EC50 = 34.4 mg/l

CONCLUSIONS

The EC50 (6hr) of the test substance was found to be 22.2 mg/l (at 680 nm) and 34.4 mg/ (at 750 nm).

DATA QUALITY

Information taken from IUCLID database.

Purity of the test substance: information not available.

GLP study: data not available

REFERENCES

IUCLID Data Sheet

4.3 Toxicity to Aquatic Plants e.g. Algae

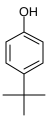
Study Ref.: Huels AG Marl

11.3 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP (Y/N): No.

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 4.1 mg/l

ChV (96hr) 0.94 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 4.1 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

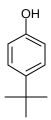
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: information not available.

METHOD

Daphnien- Kurzzeittest, DIN 38412 Teil 11, bestimmung der Wirkung von Wasserinhaltsstoffen auf Kleinkrebse

Test type: Data not available

GLP (Y/N): Data not available

Year study performed: 1982

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 48 hours

Test conditions:

Temperature: 20℃

pH: 8.0 ± 0.2

Total hardness: 2.4 mmol/l

(Ca/Mg ratio = 4:1, Na/K ratio = 10:1)

RESULTS

EC0 (48hr) 2.6 mg/l

EC50 (48hr) 3.9 mg/l

EC100 (48hr) 7.1 mg/l

CONCLUSIONS

The EC50 (48hr) of the test substance is 3.9 mg/l.

DATA QUALITY

Information taken from IUCLID database.

Purity of the test substance: information not available.

GLP study: data not available.

REFERENCES

IUCLID Data Sheet

4.2 Acute Toxicity to Aquatic Invertebrates

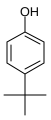
Study Ref.: Huels AG Marl

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA) - ADDITION

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: 99.5%.

METHOD

OECD TG 202

Test type: Semi-static, open-system

GLP (Y/N): No.

Year study performed: 1995

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 48 hours

20 daphnids (4replicates; 5 organisms per replicate) were exposed to nominal concentrations of 1.0, 1.8, 3.2, 5.6, and 10 mg/l. 10 mg/l DMSO and HCO-40 (4:1 weight ratio) was used as a solubiliser. 10 mg/l solubiliser and de-chlorinated tap water was used as a control.

RESULTS

EC50 (24hr) 7.3 mg/l

EC50 (48hr) 6.7 mg/l

NOEC < 1.0 mg/l

The EC50 (48 hour) was determined to be 6.7 mg/l with 95% confidence limits of 5.2 mg/l to 9.7 mg/l.

CONCLUSIONS

The EC50 (48hr) of the test substance is 6.7 mg/l.

DATA QUALITY

Taken from SIDS Dossier

REFERENCES

Environment Agency of Japan (1995) as cited in the SIDS Dossier on the HPV Phase-4 Chemical, ptert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

OTHER

In another OECD TG 202 test on *Daphnia magna*, the following results were obtained:

EC50 (24 hour) 4.2-7.3 mg/l (based on immobilisation)

EC50 (48 hour) 3.4-3.9 mg/l (based on immobilisation)

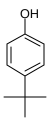
Waern (2000) and SIDS (2000) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

12.3 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA) - ADDITION

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: 99.5%.

METHOD

OECD TG 202

Test type: Static

GLP (Y/N): Unknown

Year study performed: 1989

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 48 hours

Ten test animals (6-24 h old) were exposed to 3-4 concentrations of this chemical dissolved into the dilution water (pH 8.0 +0.2, kept at 20 C, DO not below 4.0 mg/l)) in a 50 ml beaker for 48 h. Swimming daphnids were counted.

RESULTS

EC50 (24hr) 3.4 mg/l

EC50 (48hr) 3.4 mg/l

EC0 (48hr) 0.34 mg/l

The EC50 (48 hour) was determined to be 3.4 mg/l.

CONCLUSIONS

The EC50 (48hr) of the test substance is 3.4 mg/l.

DATA QUALITY

Taken from SIDS Dossier

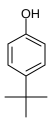
REFERENCES

Kuhn, R., Pattard, M., Pernak, K.D., and Winter, A. (1989) Water Res., 23, 495-499, as cited in the SIDS Dossier on the HPV Phase-4 Chemical, p-tert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

12.4 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

p- tert-butylphenol



CAS No. 98-54-4

IUCLID Identification Number: 98-54-4

Purity of the test substance: unknown.

METHOD

Acute Toxicity to Non-Standard Species

Test type: unknown

GLP (Y/N): No data

Year study performed: 1981

Species: Crangon septemspinosa (Shrimp)

Analytical monitoring: No data

Exposure period: 96 hours

RESULTS

LC50 (96hr) 1.9 mg/l

The LC50 (96 hour) was determined to be 1.9 mg/l.

CONCLUSIONS

The LC50 (96 hour) was determined to be 1.9 mg/l.

DATA QUALITY

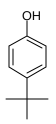
Unknown

REFERENCES

McLeese et al (1981) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

TEST SUBSTANCE

p- tert-butylphenol



CAS No. 98-54-4

IUCLID Identification Number: 98-54-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP (Y/N): No.

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 2.1 mg/l

ChV (21day) 0.32 mg/l

Remark: log Kow used 3.42 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 2.1 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

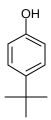
Calculation performed for this HPV submission.

12.6 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES (DAPHNIA) - ADDITION

TEST SUBSTANCE

p- tert-butylphenol

CAS No. 98-54-4



IUCLID Identification Number: 98-54-4

Purity of the test substance: 99.5%.

METHOD

OECD TG 202 (1984)

Test type: Semi-static, open-system

GLP (Y/N): No.

Year study performed: 1995

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 21 days

Endpoint: reproduction rate

40 daphnids (4 replicates; 10 daphnids per replicate) were exposed to five concentrations (0.073, 0.23, 0.73, 2.3, and 7.3 mg/l) in de-chlorinated tap water (pH: 7.6 to 8.0; Hardness 48 to 111 mg/l). DMSO and HCO-40 (4:1 mixture, 7.3 mg/l) was added as a solubiliser.

RESULTS

EC50 (21 days) 2.0 mg/l

NOEC 7.3 mg/l

NOEC 2.3 mg/l

The reproduction rate EC50 (21 days) was determined to be 2.0 mg/l.

CONCLUSIONS

The EC50 (48hr) of the test substance is 2.0 mg/l, endpoint reproduction rate.

DATA QUALITY

Taken from SIDS Dossier

REFERENCES

Environment Agency of Japan (1995) as cited in the SIDS Dossier on the HPV Phase-4 Chemical, ptert-Butylphenol, sponsor country: Japan, dated December 1, 1999.

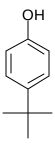
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

EEC Directive 67/548 Method B1; OECD Guidelines No. 401

GLP (Y/N): Yes.

Year study performed: 1991

Species: Sprague-Dawley rats

No. of animals per sex per dose: 10 (five male and five female),

Vehicle: Arachis oil B.P.

Route of administration: Oral gavage

Dose level: 2000 mg/kg

10 rats were given a single oral dose of test material as a solution/suspension at a dose level of 2000 mg/kg.

RESULTS

There were no deaths and no signs of systemic toxicity during the study. No abnormalities were noted at necropsy.

LD50 >2000 mg/kg

CONCLUSIONS

Essentially non-toxic. No symbol and risk phrases are required according to EEC labelling regulations.

DATA QUALITY

The study was performed to OECD guidelines and GLP.

REFERENCES

Range-finding acute oral toxicity test in the rat Project Number 47/1596, Sandoz Chemicals Test Report. Issued by Safepharm Laboratories, P.O Box No45, Derby, DE1 2BT, U.K.

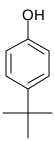
OTHER

There are data for 6 other acute oral toxicity study mentioned in the IUCLID database. These all give a value for LD_{50} of between 2990 and 5360 mg/kg.

13.2 ACUTE TOXICITY - ADDITION ORAL

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

OECD Guidelines No. 401 (1981)

GLP (Y/N): No.

Year study performed: 1985

Species: Rat

RESULTS

LD50: 4000 mg/kg b.w.

DATA QUALITY

The study was performed to OECD guidelines.

REFERENCES

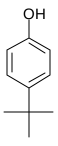
Huels-Bericht, unpublished report (1985) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

13.3 ACUTE TOXICITY - ADDITION ORAL

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

Purity: 99%

METHOD

Not specified

GLP (Y/N): No.

Year study performed: 1988

Species: Sprague-Dawley rats

Vehicle: 25% (w/v) suspension in corn oil.

Route of administration: Stomach intubation.

Dose level: 2,500, 3,500, 5,000 and 10,000 (males only) mg/kg

RESULTS

LD50 (male): 5,360 mg/kg b.w.

LD50 (female): 3,620 mg/kg b.w.

CONCLUSIONS

Signs of toxicity subsided in survivors at 3 to 7 days after dosing. Principal signs of toxicity included sluggishness, unsteady gait, prostration, unkempt appearance, and nasal discharge. Deaths were induced from 2 hours to 5 days after dosing. In rats that died during the study, there was mottling of the lungs and livers as principal macroscopic lesions in the females but no significant gross lesions in the males.

DATA QUALITY

This test was performed to OECD guidelines.

REFERENCES

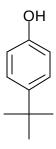
Klonne, D.R. et al. (1988) Drug Chem. Toxicol. 11, 43. as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

13.4 ACUTE TOXICITY INHALATION

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Not specified

Type: Static exposure to substantially saturated vapor

GLP (Y/N): No data

Year study performed: 1988

Species: Rats

Exposure time: 6 hours

100 g of test substance was placed in a 1201 chamber for ca. 18 hours at ambient temperature prior to the introduction of the rats.

RESULTS

No effects on body weight, clinical signs, mortality or necropsy observations were found.

CONCLUSION

No effect in rats from exposure to saturated vapor of test substance.

DATA QUALITY

Information from IUCLID database

No data for GLP

REFERENCES

IUCLID Dataset created by European COMMISSION-European Chemicals Bureau Created 19 Feb 2000

OTHER - ADDITIONAL

In an additional inhalation toxicity study, 5 male and 5 female Sprague-Dawley rats were exposed for 4 hours to p-t-butylphenol (99% purity). Exposure was conducted as dust aerosol of 5,600 mg/cm3 with an additional vapour component of 30 mg/cm3. Within one to two days following exposure, 1/5 rats of each sex died with marked dark red or purple discoloration of the lungs and kidneys. Weight

loss was observed in survivors of both sexes day 7, with an increase in weight observed on day 14. Clinical signs observed on the day of exposure up to 7 days post-exposure included signs of mucosal irritation (perinasal, perioral, and periocular encrustation) and signs of respiratory distress (audible respiration, gasping, and a decreased respiration rate).

REFERENCES - ADDITION

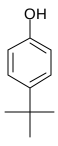
Klonne, D.R. et al. (1988) Drug Chem. Toxicol. 11, 43. as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

13.5 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

OECD Guidelines No. 404; EEC Directive 84/449 Method B4

Type: Acute Dermal Irritation/Corrosion

GLP (Y/N): Yes.

Year study performed: 1991

Species: New Zealand White rabbit

No. of animals per sex per dose: 3

Vehicle: Test substance was moistened with distilled water

Route of administration: Semi-occluded application to intact skin.

Exposure time: 4 hours

Dose level: 0.5 g

RESULTS

Severe irritant to rabbit skin according to the Draize classification scheme.

Slight to well-defined erythema was noted at all treated skin sites one hour after patch removal and persisted for at least 24 hours.

CONCLUSION

The symbol "Xi" and risk phrase R38 "irritating to skin" are therefore required.

DATA QUALITY

Study performed to OECD guidelines and GLP.

REFERENCES

Acute Dermal Irritation Test in the rabbit - Project Number: 47/1597

SafePharm Laboratories Limited, P.O Box No 45, Derby, DE1 2BT, U.K.

OTHER

A paper in Drug and Chemical Toxicology, 11(10, 43-54 (1988) entitled "Acute Toxicity and Primary Irritation of Para-Tertiary Butylphenol", examined Peroral, percutaneous inhalation toxicity and skin and eye irritation. The main conclusion from this paper was acute exposure to the test substance is associated with persistent ocular injury, and to a lesser extent, from dermal contact.

OTHER - ADDITION

In another acute dermal toxicity study, moderate irritation resulted when the skin (moistened with distilled water) of 6 New Zealand White rabbits was exposed to 500 mg of p-t-butylphenol (99% pure) for 4 hours via a Standard Draize test. The majority of the rabbits (4/6) showed no signs of skin irritation. Minor transient erythema and desquamation was evident in one of the remaining two rabbits. The second remaining rabbit exhibited slight edema, dermal necrosis, scab formation, and desquamation. The skin appeared to be normal on day 17, post-exposure.

REFERENCES - ADDITION

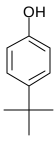
Klonne, D.R. et al. (1988) Drug Chem. Toxicol. 11, 43. as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

15.1 GENETIC TOXICITY IN VITRO (CHROMOSOMAL ABERRATIONS)

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

Purity of the test substance: > 95%.

METHOD

Dean and Hodson-Walker 1979

Type: Chromosome aberration assay

GLP (Y/N): No.

Year study performed: 1984

Species/strain: Rat liver cells

Metabolic activation: No

Exposure period: 24 hours

and all formulations were shown to be stable for at least one working day.

Positive controls: Ethyl methanesulphonate, methyl methanesulphonate and cyclophosphamide were prepared as aqueous solutions.

4-nitroquinoline, N-oxide and benzo(alpha)pyrene were dissolved in dimethyl sulphoxide.

inhibition effects noted.

was centrifuged and the cells fixed in 3 changes of fixative solutions (methanol: acetic acid: 3:1).

Chromosome preparations were made on microscopes slides and stained with Giemsa stain. The preparations were randomly coded and 100 cells from each culture were analysed microscopically.

RESULTS

There was no evidence of mutagenic activity

CONCLUSIONS

Test substance was considered to be non-mutagenic in this test system.

DATA QUALITY

Not performed to GLP.

Not a recognised test method.

No metabolic activation used.

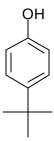
REFERENCES

Mutation Research, 153 (1985) 57-77

Genetic toxicology testing of 41 industrial chemicals.

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

Purity of the test substance: 99.9%.

METHOD

Guidelines for Screening Mutagenicity Testing of Chemicals, Japan and OECD TG 473

Type: Chromosome aberration test

GLP (Y/N): Yes.

Year study performed: 1996

Type of cells used: Chinese hamster lung (CHL/IU) cell

Test method:

Solvent: Dimethylsulfoxide

Positive control: -S9, Mitomycin C; +S9, Cyclophosphamide

Dose: -S9 (continuous treatment): 0, 0.013, 0.025, 0.05 mg/ml

-S9 (short-term treatment): 0, 0.02, 0.04, 0.08 mg/ml

+S9 (short-term treatment): 0, 0.013, 0.025, 0.05 mg/ml

S-9: Rat liver, induced with phenobarbital and 5,6-benzoflavone

Metabolic activation: With and without activation.

Plates/test: 2

RESULTS

Cytotoxicity conc: Without metabolic activation: (continuous treatment): 0.025 mg/ml

Without metabolic activation: (short-term treatment): 0.08 mg/ml

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With metabolic activation: (short-term treatment): not observed

Precipitation conc.

Genotoxic effects: clastogenicity polyploidy

With metabolic activation: positive positive Without metabolic activation: negative positive

CONCLUSIONS

Structural chromosome aberration was observed in CHL/IU cells with exogenous metabolic activation, by short-term treatment, in 6.5% - 12.0% of cells at all cell concentrations studied. This chemical also induced polyploidy with and without an exogenous metabolic activation system. The high incidences of polyploidy cell were observed after 48 hour continuous treatment but the response was apparent only at cytotoxic concentrations. Other increases of polyploidy were small incidences.

DATA QUALITY

Study performed to OECD guidelines and GLP.

REFERENCES

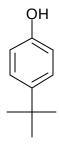
Ministry of Health and Welfare: Japan, Toxicity Testing Reports of Environmental Chemicals 4, 277-304 (1996) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

15.3 GENETIC TOXICITY IN VITRO (CHROMOSOMAL ABERRATIONS) - ADDITION

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

OECD Guidelines for Testing of Chemicals (1981) No. 473 "Genetic Toxicology: Chromosome Aberration Test" and Method B10 of Commission Directive 84/449/EEC

Type: Chromosome aberration test

GLP (Y/N): Yes.

Year study performed: 1992

Type of cells used: Rat lymphocytes

Test method:

Solvent: Dimethylsulfoxide

Positive control: -S9, Ethyl methanesulphonate at 500 µg/ml

+S9: Cyclophosphamide at 4.2 µg/ml

Dose: -S9 (20-hour cultures): $1.0 - 125 \mu g/ml$

+S9 (20-hour cultures): $1.88 - 120 \mu g/ml$

 $+S9 (30-hour culture): 1.88 - 60 \mu g/ml$

S-9: Rat liver, induced with Aroclor 1254

Metabolic activation: With and without activation.

Plates/test: 2

RESULTS

Cytotoxicity conc: Without metabolic activation: 60.0 µg/ml and more dose

(20- and 30-hour cultures)

Without metabolic activation: 30.0 µg/ml and more dose

Precipitation conc.

Genotoxic effects: clastogenicity polyploidy

With metabolic activation: negative negative without metabolic activation: negative negative

CONCLUSIONS

This test substance is not considered to be mutagenic in this test system.

DATA QUALITY

Performed to GLP. Except that the concentration, homogeneity and stability of test material preparations were not determined by analysis and that this specific study may not have been subject to procedure inspection by the Quality Assurance Unit.

REFERENCES

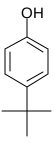
Safepharm Laboratories Ltd., on behalf of DOW Europe S.A., Project No. 44/903, unpublished (1992b) as cited in the SIDS Dossier on the HPV Phase 4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

15.4 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY)

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

Purity of the test substance: > 95%

METHOD

Method: Ames et al (1975)

Type: Reverse mutation assay, Bacterial - Plate incorporation and pre-incubation methods used

GLP (Y/N): No.

Year study performed: 1980

Species/strain: Salmonella typhimurium TA 1535, TA 1537, TA 1538, TA 98 and TA 100 and E.coli WP2 and WP2uvrA

Metabolic activation: S-9 mix (with and without).

Plate incorporation assay

cytotoxicity assay has been carried out to assess both the cytotoxicity of the test compound and its solubility in the top agar. The amounts to be used in the mutation assays were selected on this basis. 0.1 ml of a dilution (1: 20000) of an overnight bacterial culture was added to 2 ml top agar, together with $20 \mu l$ test compound to give the following final amounts:

Dose/concentration without S -9 mix: 125, 250, 500, 1000, 2000 and 4000 µg/plate

Dose/concentration with S-9 mix: 125, 250, 500, 1000, 2000 and 4000 μg/plate

pyrene, 7,12 dimethylbenzanthracene and 4-nitroquinoline N-oxide were dissolved in dimethylsulphoxide (DMSO).

Pre-incubation assay

The method used was that described by Brooks and Dean (1981). Bacteria (0.5 ml) with and without the S-9 mix wee incubated at 37°C with the test solution (0.1 ml) or solvent for 30 minutes before incorporation of 0.5 ml of this pre-incubation mixture into 2 ml of top agar. All assays were carried out at least in triplicate.

The stability of the formulation material and positive control compounds were assessed by appropriate analytical methods and formulations were shown to be stable for at least one working day.

RESULTS

The test substance did not cause any mutations in either test system.

CONCLUSIONS

Test substance is considered to be non-mutagenic.

DATA QUALITY

Not performed to GLP, but a recognised test method.

REFERENCES

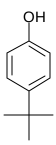
Mutation Research, 153 (1985) 57-77.

Genetic toxicology testing of 41 industrial chemicals.

15.5 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY) - ADDITION

TEST SUBSTANCE

CAS No. 98-54-4



IUCLID ID Number 98-54-4

Purity of the test substance: 99.9%

METHOD

Guidelines for Screening Mutagenicity Testing of Chemicals Japan, and OECD TG (471 and 472)

Type: Bacterial gene mutation assay.

GLP (Y/N): Yes.

Year study performed: 1996

System of testing: S. typhimurium TA98, TA100, TA1535, TA1537

E.coli WP2 uvrA

Dose: -S9 mix: 0, 15.6, 31.3, 62.5, 125, 250 and 500 μg/plate

(TA100, TA1535, TA98, TA1537), 0, 31.3 – 1000 μg/plate (WP2)

+S9 mix: Same as -S9 mix.

Metabolic activation: With and without activation.

S9: Rat liver, induced with phenobarbital and 5,6-benzoflavone

Plate/test: 3

Number of replicates: 2

RESULTS

Cytotoxicity conc: With metabolic activation: 500 µg/plate (five strains)

Without metabolic activation: 500 µg/plate (TA100, TA1535, TA 1537)

1000 µg/plate (WP2 and TA98)

Precipitation conc.

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With metabolic activation: negative

Without metabolic activation: negative

CONCLUSIONS

This test substance is not considered to be mutagenic under these test conditions.

DATA QUALITY

The study was performed to OECD guidelines and GLP.

REFERENCES

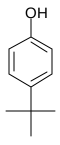
Ministry of Health and Welfare: Japan, Toxicity Testing Reports of Environmental Chemicals 4, 277-304 (1996) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

15.6 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY) - ADDITION

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Guidelines for bacterial mutagenicity testing published by the major Japanese Regulatory Authorities including MITI, MHW, MOL and MAFF

Type: Bacterial gene mutation assay

GLP (Y/N): Yes.

Year study performed: 1992

System of testing: S. typhimurium TA1535, TA1537, TA98, TA100

E.coli WP2 uvrA

Test method:

Solvent: Dimethylsulfoxide

Dose: 1.6, 8, 40, 200, 1000 μg/ml

Metabolic activation: With and without activation.

RESULTS

Cytotoxicity conc: With metabolic activation: 1000 µg/ml (all strains)

Without metabolic activation: 1000 µg/ml (all strains)

Precipitation conc.

Genotoxic effects:

With metabolic activation: negative

Without metabolic activation: negative

Positive control: With metabolic activation:

2-Aminoanthracene at 2 μ g/plate for TA 1535 and 10 μ g/plate for WP2 uvrA

Benzo(a)pyrene at 5 µg/plate for TA100, TA 1537 and TA98

Without metabolic activation:

N-ethyl-N'nito-N-nitrosoguanidine at 2 μ g/plate for WP2 uvrA, 3 μ g/plate for TA100 and 8 μ g/plate for TA 1535

9-Aminoacridine at 80 µg/plate for TA1537

4-Nitroquinoline-1-oxide at 0.2 μg/plate for TA98

CONCLUSIONS

The test material is considered to be non-mutagenic in this test system.

DATA QUALITY

Performed to GLP. Except that the concentration, homogeneity and stability of test material preparations were not determined by analysis and that this specific study may not have been subject to procedure inspection by the Quality Assurance Unit.

REFERENCES

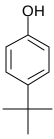
Safepharm Laboratories Ltd., on behalf of DOW Europe S.A., Project No. 44/901, unpublished (1992a) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

15.7 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY)

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Method: Honma et al 1998

Type: Gene mutation assay- Mouse lymphoma

GLP (Y/N): No.

Year study performed: 1999

Species/strain: L5178y tk+/- clone, mouse lymphoma cells

Metabolic activation: No

Concentrations tested: Dose/concentration without S-9 mix: 10, 20, 30, 40, 50, 60, 70 and 80 µg/ml

Dose/concentration with S-9 mix: N/A

Exposure time: 3 and 24 hours. This study was undertaken to validate the mouse lymphoma assay as an alternative to the chromosome assay, and therefore included a later harvest time.



positive responses in the 24 hour treatment mouse lymphoma assay. Combining results of the previous and the current studies, 31 of 34 chromosome aberration positive chemicals exhibited positive responses in the mouse lymphoma assay, suggesting that the sensitivity of the MLA with 24 hour treatment approaches that of the chromosome aberration test.

*M. Honma

RESULTS

No evidence of mutagenic response after 3 hours treatment (normal exposure time). Although positive response seen at 24 hour treatment this data is not relevant as only used as a demonstration:

Following the advice from the ICH, a collaborative study of the mouse lymphoma assay (MLA) was conducted. The major aim of the study was to validate the MLA as an alternative to the chromosome aberration test (CA). 34 CA-positive chemicals, mostly clastogens and spindle poisons, were tested and investigated for the positive response in the MLA. Nine negative and five marginal responses were obtained, indicating that at least 26% (9/34) of clastogens were not detected by the MLA. Therefore, it was concluded that the MLA is not equivalent to the CA and could not be regarded as an alternative to the CA test. The discordant results between MLA and the CA however, might have been due to differing treatment periods (the standard MLA employs short 3 or 4 hr treatment) while CA employs long-term (24 or 48 hr) as well as short-term (4 or 6 hr) treatments. The short-term MLA maybe unable to detect some clastogens and spindle poisons. In this study, a long-term MLA was used to reexamine 15 chemicals that were judged to be negative or inconclusive in the collaborative MLA study.

In a previous collaborative study 34 chemicals (positive in the CA but negative in the bacterial reverse mutation assay) were examined. Fourteen could not be judged to be positive in the MLA. This collaborative study, however, suggested that the MLA may not have detected some clastogens and spindle poisons because the treatment period (3hr) may have been too short. Among the fourteen chemicals, four did not produce the target toxicity even at the maximum dose tested (5 mg/ml). Furthermore, two chemicals were positive in the CA with 24 hr treatment but not with pulse treatment (6 hr) indicating that longer exposure was needed for expression of their genotoxicity.

The MLA generally employs 3 or 4 hr treatment. Extended treatment times however are considered inappropriate in the MLA because prolonged toxicity delays the cell cycle, which in turn prolongs the time for suitable mutation fixation and thus results in fluctuations in mutation frequencies. Long term treatment was, however, sometimes employed in other mammalian cell gene mutation systems such as CHO/HPRT and TK6/TK. However, even with long term treatment the complete concordance between the CA and MLA would not be expected; the endpoints are different and different biological processes might affect the results. Thus in order to compare the inter-changability of the CA and MLA, the effectiveness of the long term treatment in the MLA needs to be evaluated.

CONCLUSIONS

Test substance is considered to be non-mutagenic after the normal 3 hour treatment time in the absence of S9 mix.

DATA OUALITY

Not performed to GLP and not tested in the presence of metabolic activation.

REFERENCES

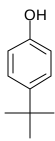
Mutagenesis vol. 14 no 1 pp 23-29, 1999

15.8 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY) - ADDITION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Type: Mutation of L5178Y mouse lymphoma cells at the thymidine kinase TK+/- locus. Fluctuation assay.

GLP (Y/N): Yes.

Year study performed: 1992

Type of cells used: L5178Y mouse lymphoma cells (heterozygous at the thymidine kinase locus)

Test method:

Solvent: Dimethylsulfoxide

Positive control: -S9: Ethyl methanesulphonate at 931.5 µg/ml

+S9: Cyclophosphamide at 5 µg/ml

Dose: 5.0, 10.0, 20, 40, 80 μg/ml

S-9: Rat liver, induced with Aroclor 1254

Metabolic activation: With and without activation.

Plates/test: 2

RESULTS

Cytotoxicity conc: 80 µg/ml

Precipitation conc.

Genotoxic effects:

With metabolic activation: negative

Without metabolic activation: negative

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CONCLUSIONS

This test material is not considered to be mutagenic in this test system.

DATA QUALITY

Performed to GLP. Except that the concentration, homogeneity and stability of test material preparations were not determined by analysis and that this specific study may not have been subject to procedure inspection by the Quality Assurance Unit.

REFERENCES

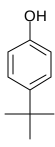
Safepharm Laboratories Ltd., on behalf of DOW Europe S.A., Project No. 44/902, unpublished (1992c) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

16.1 REPEATED DOSE TOXICITY 20 WEEKS DIETARY

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



METHOD

Study design: 20 week dietary study, non-Guideline, on 13 phenolic compounds

GLP (Y/N): No information

Year study performed: 1986

Species and sex: Male Syrian Golden Hamster, 6 weeks of age at start, in groups of 15

Route of administration: Oral, by incorporation in feed

Exposure period: Continuous for 20 weeks

Dose: 15 000 ppm. Dosage was based on one quarter of the LD50 value determined for rats.

Liver and kidneys weighed, preserved with cheek pouch, stomach, oesophagus, lung, pancreas and urinary bladder. These tissues examined histologically. Autoradiography performed for stomach and bladder tissue, labelling increased DNA synthesis with ³H-thymidine and thus identifying dividing cells.

RESULTS (for PTBP)

Average body weight at the end of the study was 5% less than that of the control group. Average liver weight (relative to body weight) was increased by 21%.

Prominent thickening of the forestomach epithelium with a keratin-like white substance observed in the posterior and anterior walls along the lesser curvature and adjacent to the oesophagus.

Chemical	No. of Body wt hamsters (g)	Liver wt. (g/100g	No. of hamsters (%) with:				
		(g)	body wt)	Mild hyperplasia	Moderate hyperplasia	Severe hyperplasia	Papillomato us lesions
PTBP	15	192 ± 14	4.6± 0.4	15 (100) ***	12 (80) ***	11 (73.3)***	7 (46.7) ***
Basal diet	15	203 ± 23	3.8 ± 0.4	7 (46.7)	1 (6.7)	0	0

***P < 0.001; ** P < 0.01; *P < 0.05

No abnormal findings were observed in liver, kidneys, cheek pouch, lung, pancreas and urinary bladder.

Chemical	No. of hamsters	Labelling index (Mean \pm SD of labelled cells/100 cells)			
		Forestomach	Pyloric region	Urinary bladder	
PTBP	3	34.3 ± 6.4 **	10.7 ± 0.4	0.13 ± 0.06	
Basal diet	3	12.5 ± 4.7	7.3 ± 1.9	0.08 ± 0.14	

^{**} P < 0.01; *P< 0.05

CONCLUSION

PTBP may have carcinogenic effects on the forestomach of hamsters. The significance of the findings (in a structure that does not occur in humans) for human cancer hazard is not known.

DATA QUALITY

No information to confirm if this was a GLP study.

REFERENCE

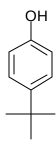
Hirose, M. et al. (1986) Carcinogenesis 7, 1285-1289

16.2 REPEATED DOSE TOXICITY – ORAL – ADDITION 14 DAY – DOSE FINDING

TEST SUBSTANCE

p-tert-Butylphenol

CAS No. 98-54-4



Purity of substance tested: 99.9%

METHOD

Dose finding study

GLP (Y/N): No.

Year study performed: unknown

Species and sex: Male/Female Rats/Crj: CD (SD)

Route of administration: Oral gavage

Exposure period: 14 days, daily

Post-exposure Observation Period: 1 day

Dose: 0 (Vehicle), 250, 500, 1000 mg/kg/day

Control group: Yes; 0.5% aqueous methyl cellulose

LOEL: 250 mg/kg/day

RESULTS

At 1000 mg/kg, the body weights were 10% lower than the control at day 9, and 3/5 females and 1/5 males died up to day 9. At this time, all other survivors were dissected but no toxic sign was observed by necropsy. Two females continuously showed difficulty of breathing with noisy respiratory sound.

CONCLUSION

There was no systemic toxicity of this chemical but the tolerable dose of this chemical would be around 250 mg/kg/day for OECD Combined Repeat Dose and Reproductive/Developmental Toxicity Screening Test, because of difficulty of breathing, probably caused by irritation.

DATA QUALITY

Not performed to GLP

REFERENCE

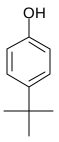
Ministry of Health and Welfare, Japan: Unpublished, as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

16.3 REPEATED DOSE TOXICITY – ORAL - ADDITION (44 DAYS – MALE), (14 DAYS PRE-MATE TO DAY 4 LACTATION – FEMALE)

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



Purity: 99.9 %

METHOD

OECD Combined Repeat Dose and Reproductive/Developmental Toxicity Screening Test (OECD TG 422)

GLP (Y/N): Yes.

Year study performed: 1996

Species and sex: Male/Female Rats/Crj: CD (SD)

Route of administration: Oral gavage

Exposure period: Male: 44 days

Female: from 14 days before mating to day 4 of lactation

Dose: 0 (Vehicle), 20, 60, 200 mg/kg/day

Control group: Yes; 0.5% aqueous methyl cellulose

RESULTS

NOAEL: 200 mg/kg/day as systemic toxicity

No treatment related changes were observed except noisy respiratory sound in a few females of the 200 mg/kg group. Only plasma albumin in the 200 mg/kg males was decreased.

CONCLUSION

NOAEL: 200 mg/kg/day as systemic toxicity

Noisy respiration sound was likely related to irritation of the respiratory tract caused by the oral administration of the chemical.

DATA QUALITY

Study performed to OECD guidelines and GLP.

REFERENCE

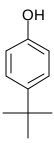
Ministry of Health and Welfare: Japan, Toxicity Testing Reports of Environmental Chemicals 4, 277-304 (1996) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

16.4 REPEATED DOSE TOXICITY 51 WEEKS CARCINOGENICITY DIETARY

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Method: Other, not specified

Type: 51 weeks

GLP (Y/N): No data

Year study performed: 1988

Species and sex: Male Fischer 344 rats

Route of admin: Oral feed

Exposure period: 51 weeks

Post observation period: 1 week

Doses: 15000 ppm

Control group: Yes, concurrent no treatment

RESULTS

Carcinogenicity: Average body weights and relative weights were decreased significantly (p<0.001): relative kidney weights were significantly increased; histological changes in the forestomach.

Hyperplasia: 14/15 rats, control 0/10

Papilloma: 1/15, control 0/10

Carcinoma within hyperplastic epithelium without invasion of the submucosa (carcinoma "in situ") 0/15, control 0/15

Squamous cell carcinoma: 0/15, control 0/15

No hyperplasia or carcinoma found in the glandular stomach, and no tumours seen in oesophagus, liver kidneys or intestines.

Modifying effect of 4-tert-butylphenol on N-Methyl-N''-nitro-N-nitrosoguanidine (MNNG)- induced forestomach and glandular stomach carcinogenesis:

At 6 weeks of age, the animals were given 150 mg/kg bw MNNG in saline by stomach tube; one week later rats were treated. Animals, which were treated with MNNG, followed by basal diet served as controls.

Average body weight decreased significantly, and relative liver and kidney weights increased significantly.

Histological changes in the forestomach:

Hyperplasia: 20/20 rats, control 19/19

Papilloma: 19/29 rats, control 13/19

Carcinoma "in situ": 8/20, control 11/19

Squamous cell carcinoma: 15/20 rats (p<0.01) control 5/19

Leiomyosarcoma induced in one rat: no tumours seen in other organs.

CONCLUSION

Test substance has a very weak or lacks carcinogenic activity, but enhances forestomach carcinogenesis in rats treated with MNNG.

DATA QUALITY

Information taken from IUCLID database. "The data in this summary is supportive rather than key information".

No data to confirm if GLP

REFERENCES

Huels Ag Marl

Hirose, M et al (1988): Cancer Res. 48, 5310-5315

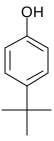
IUCLID Dataset created by European COMMISSION-European Chemicals Bureau Created 19 Feb 2000

17.1 TOXICITY TO REPRODUCTION (ESTROGENIC ACTIVITY) - ADDITION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



IUCLID ID Number 98-54-4

METHOD

Guideline followed: Estrogenic Activity Yeast Screen

GLP (Y/N): No.

Year study performed: 1997

Yeast is transfected with two plasmids, one carrying human estrogen responsive element and reporter gene Lac-Z. The estrogenic response is measured and compared to that of 17ß-estradiol.

RESULTS

In yeast transfected with two plasmids, one carrying human estrogen responsive element and reporter gene Lac-Z, p-t-butylphenol indicated very weakly estrogenic activity. The estrogenic transcriptional activity of p-t-butylphenol was 1,500,000 fold less potent than 17ß-estradiol.

CONCLUSION

The estrogenic transcriptional activity of p-t-butylphenol was 1,500,000 fold less potent than 17ß-estradiol.

DATA QUALITY

Included in the SIDS Dossier.

Not GLP or OECD compliant.

REFERENCES

Routledge E.J., and Sumpter, J.P. (1997) Estrogenic Activity of Surfactants and some of their degradation products assessed using a re-combinant yeast screen. J. Biol. Chem., 272:3280, as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

OTHER

E Screen Relative Potency = 0.0003

Yeast Screen Relative Potency = 0.2

Potency Relative to estradiol = 100

REFERENCE - OTHER

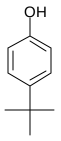
Waern (2000) as summarised in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

17.2 TOXICITY TO REPRODUCTION - ADDITION

TEST SUBSTANCE

*p-tert-*Butylphenol

CAS No. 98-54-4



Purity: 99.9 %

METHOD

OECD Combined Repeat Dose and Reproductive/Developmental Toxicity Screening Test (OECD TG 422)

GLP (Y/N): Yes.

Year study performed: 1996

Species and sex: Male/Female Rats/Crj: CD (SD)

Route of administration: Oral gavage

Exposure period: Male: 44 days

Female: from 14 days before mating to day 4 of lactation

Dose: 0 (Vehicle), 20, 60, 200 mg/kg/day

Control group: Yes; 0.5% aqueous methyl cellulose

RESULTS

NOAEL: 200 mg/kg/day as systemic toxicity

No treatment related changes were observed except noisy respiratory sound in a few females of the 200 mg/kg group. Only plasma albumin in the 200 mg/kg males was decreased.

There were no treatment related toxic effects on pregnant and lactating females or their offspring.

NOAEL: 200 mg/kg/day as systemic toxicity

No treatment related toxic effects on pregnant and lactating females or their offspring were seen with this chemical in this test system.

DATA QUALITY

Study performed to OECD guidelines and GLP.

REFERENCE

Ministry of Health and Welfare: Japan, Toxicity Testing Reports of Environmental Chemicals 4, 277-304 (1996) as cited in the SIDS Dossier on the HPV Phase-4 Chemical p-tert-butylphenol, Cas # 98-54-4. Sponsor country: Japan, December 1,1999.

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: information not available in reference.

GLP (Y/N): No.

Year study performed: not known.

RESULTS

Melting point: 94 - 95°C.

CONCLUSIONS

The test substance has a melting point range of 94 - 95°C.

DATA QUALITY

Not a GLP study.

Information taken from a standard reference book.

Purity of the test substance / decomposition: information not available in reference.

REFERENCES

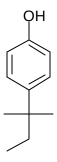
The Merck Index – Encyclopedia of Chemicals, Drugs and Biologicals. Rahway, NJ: Merck and Co., Inc., 1989, p. 1132.

2.1 BOILING POINT

TEST SUBSTANCE

*p-tert-*Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: information not available in reference.

GLP (Y/N): No.

Year study performed: not known.

RESULTS

Boiling point: 262.5°C (atmospheric pressure assumed).

CONCLUSIONS

The test substance has a boiling point of 262.5°C.

DATA QUALITY

Not a GLP study.

Information taken from a standard reference book.

Purity of the test substance / decomposition: information not available in reference.

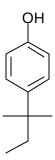
REFERENCES

The Merck Index – Encyclopedia of Chemicals, Drugs and Biologicals. Rahway, NJ: Merck and Co., Inc., 1989, p. 1132.

3.1 VAPOR PRESSURE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: calculation.

GLP (Y/N): No.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 0.00783 mmHg @ 25°C (1.04 Pa).

CONCLUSIONS

The test substance has a calculated vapor pressure of 0.00783 mmHg @ 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1

OTHER

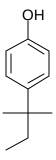
Value obtained using Modified Grain Method reported.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: information not available.

GLP (Y/N): No.

Year study performed: 1987.

RESULTS

Log Kow: 4.03.

CONCLUSIONS

The test substance has a log Kow of 4.03.

DATA QUALITY

Not a GLP study.

Information taken from literature search covering appropriate databases.

Purity of the test substance: information not available.

REFERENCES

Schultz TW; Ecotoxicology and Environmental Safety 14: 178-83 (1987)

OTHER

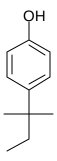
p-tert-Amylphenol has a pKa of 10.43 (Schultz TW; Ecotoxicology and Environmental Safety 14: 178 – 83 (1987).)

5.1 WATER SOLUBILITY

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No

Year study performed: not known.

RESULTS

Water solubility: 168 mg/l at 25°C.

CONCLUSIONS

The test substance has a water solubility of 168 mg/l at 25 °C.

DATA QUALITY

Not a GLP study.

Information taken from a database.

Purity of the test substance: information not available in database.

REFERENCES

Yalkowsky SH, Dannenfelser RM; Aquasol Database of Aqueous Solubility. 5th ed. Tucson, AZ Univ. of Arizona, College of Pharmacy (1992).

OTHER

p-tert-Amylphenol has a pKa of 10.43 (Schultz TW; Ecotoxicology and Environmental Safety 14: 178 – 83 (1987).)

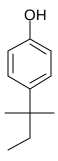
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): No.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $41.8 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 3.07 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 3.07 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

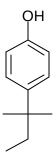
No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

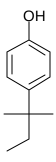
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 164
Data temperature (°C): 25
Water solubility (mg/l): 168
Vapor pressure (Pa): 1.04
Log Kow: 4.03
Melting point (°C): 94.5

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	1.88
Soil	87.0
Water	9.16
suspended sediment	0.060
Fish	0.0049
Sediment	1.93

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

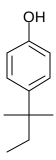
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

*p-tert-*Amylphenol

CAS No. 80-46-6



METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): No.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days/weeks

Ultimate biodegradation in weeks/months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

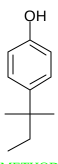
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p- tert- Amylphenol

CAS No. 80-46-6



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP (Y/N): No.

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 1.6 mg/l

ChV (30 day) 0.24 mg/l

ChV (90 day) 0.027 mg/l

Remark: log Kow used 3.91 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 1.6 mg/l.

DATA QUALITY

Estimation

ECOSAR v.0.99e

OTHER

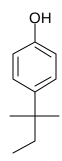
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- tert- Amylphenol

CAS No. 80-46-6



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 1.7 mg/l

ChV (96hr) 0.50 mg/l

Remark: log Kow used 3.91 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 1.7 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

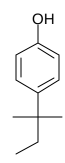
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

p- tert- Amylphenol

CAS No. 80-46-6



METHOD

Acute Toxicity to Non-Standard Species

Test type: unknown

GLP: No data

Year study performed: 1981

Species: Crangon septemspinosa (Shrimp)

Analytical monitoring: No data

Exposure period: 96 hours

RESULTS

LC50 (96hr) 1.7 mg/l

The LC50 (96 hour) was determined to be 1.7 mg/l.

CONCLUSIONS

The LC50 (96 hour) was determined to be 1.7 mg/l.

DATA QUALITY

Unknown

REFERENCES

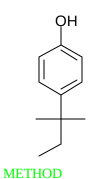
McLeese et al (1981) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- tert- Amylphenol

CAS No. 80-46-6



ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 1.5 mg/l

ChV (21day) 0.17mg/l

Remark: log Kow used 3.91 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 1.5 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

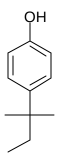
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



METHOD

Lethal dose, 50% kill

GLP: No

Year study performed: Not available (publication date was 1967)

Species: Rat.

Route of Administration: Oral.

Remarks: Details of toxic effects not reported other than lethal dose value.

RESULTS

LD50 = 1830 mg/kg.

CONCLUSIONS

LD50 = 1830 mg/kg.

DATA QUALITY

No information on guideline followed and not conducted to GLP.

Data taken from report.

REFERENCES

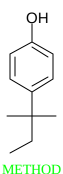
IHFCAY Industrial Hygiene Foundation of America, Chemical and Toxicological series, Bulletin. (Pittsburgh, PA) 1947-69. Volume (issue)/page/year: 6,1,1967

13.2 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

p-tert-Amylphenol

CAS No. 80-46-6



OECD Guideline No. 404

GLP (Y)

Year study performed: 1991

Species: New Zealand white Rabbit.

Number of animals: 3 + 3.

0.5 g of the test material moistened with distilled water was introduced in a semi-occluded application to the intact rabbit skin for exposure periods of 3 minutes and 4 hours.

RESULTS

A single 4-hour application produced corrosive effects at one treated skin site. These reactions included green-colored dermal necrosis, severe oedema and scabbing. Scar tissue was noted at the 14-day observation. Well-defined moderate to severe erythema and slight oedema were produced at the other two treated sites.

A single 3-minute application of the test material to the intact skin produced very slight erythema. The skin appeared normal 48 hours after treatment.

CONCLUSION

The test material is classified as corrosive according to EEC labelling regulations. The symbol C and risk phrase R34 "causes burns" is therefore required.

DATA QUALITY

The study was performed following OECD guidelines and GLP.

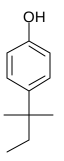
REFERENCES

Phenol, 4(1,1-Dimethylpropyl)-Acute Dermal Irritation Test in the Rabbit –Project Number 47/2098 SafePharm Laboratories Limited, P.O Box No. 45, Derby, DE1 2BT, U.K.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

*p-tert-*Amylphenol



CAS No. 80-46-6

METHOD

Method: Ames, et al., 1975; Bridges, 1972: Green and Muriel, 1976: Maron and Ames, 1983: Maron et al., 1981.

Type: Reverse mutation assay

System of testing: Bacterial: Pre incubation assay

GLP (Yes)

Year: 1993

Species/strain: Salmonella typhimurium TA 1535, TA 1537, TA 1538, TA 98 and TA 100 and E.coli WP₂ uvrA.

Metabolic activation: S9 mix (with and without) Aroclor 1254-induced male Sprague-Dawley rat liver homogenate.

Concentrations tested:

 $0.05,\,0.0167,\,0.5,\,1.67,\,5.00,\,16.7$ and $50.0\,\mu g/p$ late with and without S9 mix TEST 1 0.05, 0.0167, 0.5, 1.67, 5.00, 16.7 and 50.0 $\mu g/p$ late without S9 mix TEST 2 0.50, 0.167, 0.5, 1.67, 5.0,16.7, 50.5 167 and 500 $\mu g/p$ late with S9 mix TEST 2 0.167, 0.50, 1.67, 5.00, 16.7, 50.5 and 100 $\mu g/p$ late with S9 retest (due to cytotoxicity)

Triplicate plates

RESULTS

The test substance did not cause any increase in reverent frequency in any of the bacterial strains compared to the solvent controls.

CONCLUSIONS

Test substance is considered to be non mutagenic in this *in-vitro* test system.

DATA QUALITY

The study was performed following recognised guidelines and GLP.

REFERENCES

Pharmakon USA P.O. Box 609, Waverly, Pennsylvania 18471-0609

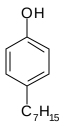
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Melting point: 73.39℃.

CONCLUSIONS

The test substance has a calculated melting point of 73.39°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30.

OTHER

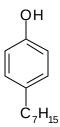
Weighted value used.

2.1 BOILING POINT

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Boiling point: 296.48°C.

CONCLUSIONS

The test substance has a calculated boiling point of 296.48°C.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30.

OTHER

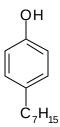
Adapted Stein & Brown method used.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 0.000279 mmHg @ 25°C (0.037 Pa).

CONCLUSIONS

The test substance has a calculated vapor pressure of 0.000279 mmHg @ 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30.

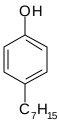
OTHER

4.1.PARTITION COEFFICIENT

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 5.01

CONCLUSIONS

The test substance has a calculated log Kow of 5.01.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63 estimate

OTHER

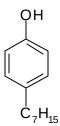
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1. WATER SOLUBILITY

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 9.645 mg/l @ 25℃.

CONCLUSIONS

The test substance has a calculated water solubility of 9.645 mg/l @ 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Log Kow value of 5.01 used (see 4.1, Partition Coefficient of *p*-heptylphenol).

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

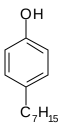
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $48.9 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.63 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.63 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

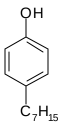
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 192
Data temperature (°C): 25
Water solubility (mg/l): 9.65
Vapor pressure (Pa): 0.037
Log Kow: 5.01
Melting point (°C): 73.39

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	0.16
Soil	96.6
Water	1.07
suspended sediment	0.067
fish	0.0055
Sediment	2.15

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

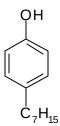
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIO DEGRADATION

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days

Ultimate biodegradation in weeks.

This material is considered to be inherently biodegradable, and would aerobically biodegrade.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

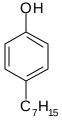
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



Purity of the test substance: information not available.

METHOD

OECD Guidelines No. 203, Fish Acute Toxicity Test.

EPA TSCA 40 CFR Part 797.1400

Test type: Static test

GLP: (Y)

Year study performed: 1997

Species: Oncorhynchus mykiss (Trout)

Test substance preparation: Water used for acclimation of the test organisms and all toxicity testing was reconstituted hard water prepared according to EPA 1993 specifications.

Analytical monitoring: none. Nominal concentrations used.

The test was conducted as a screening assay. Each treatment utilised a single replicate with 10 organisms equally distributed among each of five test treatment and diluent control.

Control:

TOC (0hr) 0 mg/l

TOC (24hr) 0.94 mg/l

Temperature $14.3 - 15.3^{\circ}$ C

Dissolved oxygen 96 – 105% saturation

pH 8.01 – 8.12

Hardness (CaCO₃) 188 mg/l

Alkalinity (CaCO₃) 114 mg/l

Total solids 404 mg/l

Test treatment:

TOC (0hr) 12.01 – 37.09 mg/l

TOC (24hr) 0.94 - 44.87 mg/l

Temperature $14.0 - 15.3^{\circ}$ C

Dissolved oxygen 93 – 106% saturation

pH 8.00 - 8.16

Hardness ($CaCO_3$) 184 – 188 mg/l

Alkalinity (CaCO₃) 108 – 114 mg/l

Total solids not applicable

There was no renewal of the test solution during the exposure period.

Exposure period: 96 hours

Statistical methods: LC50 and EC50 values were computed using a program developed by EPA (Stephen 1983) which calculates LC50 and EC50 values and their confidence limits using the probit, binomial and Spearman-Karber methods. The no observed effect concentration (NOEC) is the highest concentration of the test substance that allows at least 90% survival of exposed organisms and does not cause sub-lethal effects. This definition of the NOEC differs slightly from the TSCA test guidelines.

Deviations from protocol:

- 1.) Hourly temperatures recorded by the data logger between hours 55 and 91 of the test indicated that during the period 33 temperature reading were 16.08°C and one was 16.2°C (limit 16°C).
- 2.) Hardness of the laboratory control water was 188 mg/l CaCO_3 (protocol range $165 185 \text{ mg/l CaCO}_3$).

RESULTS

Nominal concentrations: 0, 0.10, 0.35, 3.32, 33.01 and 330.00 mg/l

Remarks: Review of solubility observations showed varying levels of product remaining on the water surface. At 0 hours, no product was observed in the 0.10, 0.35 and 3.32 mg/l concentrations. An oily sheen was observed on the water surface in the 33.01 mg/l and the entire water surface in the 330.0 mg/l concentration was covered with an oily sheen.

LC50 and EC50 values with 95% confidence limits and no-observed effect concentrations, based on nominal concentrations:

LC50 + EC50 (24hr) 1.02 mg/l (0.86 – 1.21 mg/l)

LC50 + EC50 (48hr) 1.02 mg/l (0.86 - 1.21 mg/l)

LC50 + EC50 (72hr) 0.94 mg/l (No estimate available)

LC50 + EC50 (96hr) 0.85 mg/l (0.35 - 3.32 mg/l)

NOEC (24hr) 0.35 mg/l

NOEC (48hr) 0.35 mg/l

NOEC (72hr) <0.10 mg/l

NOEC (96hr) < 0.10 mg/l

Nominal concentration	ninal concentration % Mortality			
(mg/l)	Hours	of Expo	sure	
	24	48	72	96
Control	0	0	0	0
(Level 1) 0.10	10	10	40	40
(Level 2) 0.35	0	0	0	10
(Level 3) 3.32	100	100	100	100
(Level 4) 33.01	100	100	100	100
(Level 5) 333.00	100	100	100	100

Sublethal effects were noted in the 33.01 and 330.00 mg/l concentrations as soon as the fish were added to the test solution. Fish in the 330.00 mg/l concentration were dead 5 minutes after being added to the aquarium and fish in the 33.01 mg/l concentration were dead within 25 minutes after being added to the aquarium. Fish in the 3.32 mg/l concentration exhibited sublethal effects within 20 minutes of being added to the test vessel and were dead within 3.5 hours. Sublethal effects were limited to the start of the test and no sublethal effects were observed at the 24-hour exposure.

CONCLUSIONS

Exposure of test organisms to the test substance resulted in a 96 hour LC50 of 0.85 mg/l.

The NOEC of the test substance was less than 0.10 mg/l

DATA QUALITY

Method follows OECD guideline

Information taken from the test report.

Purity of the test substance: information not available.

GLP study.

REFERENCES

Static Acute Toxicity of Lubricant additive OS#43247K to the Trout, *Oncorhynchus mykiss* Report Number 540 Sponsor – Lubrizol Corporation, 29400 Lakeland Boulevard, Wickliffe, Ohio 44092,

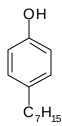
USA. Testing Facility – Envirosystems Incorporated, P.O. Box 778, 1 Lafayette Road, Hampton, New Hampshire 03842, USA.

10.2 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.40 mg/l

ChV (30 day) 0.056 mg/l

ChV (90 day) 0.010 mg/l

Remark: log Kow used 5.01 (calculated value)

water solubility 9.65 mg/l (measured)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.40 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

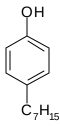
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



The test substance was assumed to be 100% active ingredient and to be stable under conditions of administration.

METHOD

OECD Guideline No. 201, Algal Growth Inhibition Test. 1984.

U.S. EPA The *Selenastrum capricornutum* Printz Algal Assay Bottle Test EPA 600/9-78-018. Environmental Research Laboratory, Corvallis, Oregon. 1978

EPA TSCA 40 CFR Part 797.1050

Test type: Static test

GLP: (Y)

Year study performed: 1998

Species: Green algae – Selenastrum capricornutum Printz

Exposure period: 96 hours, from 6th to 10th November 1997.

Analytical procedures: Nominal concentrations used.

Test substance preparation: Each of the five WAFs were prepared by stirring the mixtures of the substance and dilution water for approximately 20 hours, settling the mixtures for four hours and siphoning off the water phase containing the WAF.

The test vessels were incubated for 96 hours at $24 \pm 1^{\circ}$ C under a 24 hour light and 0 hour dark photoperiod automatically maintained with "cool-white" fluorescent lights. The dilution water was sterile enriched media adjusted to a pH of 7.5 ± 0.1 . Light intensity was maintained at 360 - 370 ft-c. The vessels were randomly arranged on a rotary shaker at the speed of 100 rpm. Algae were distributed between two replicates of each treatment at the rate of approximately 10,000 cells/ml.

Statistical methods:

Results of the toxicity test were interpreted by standard statistical techniques (Stephan C. E. 1983) when warranted. All calculations were performed using nominal concentrations of the test substance with the number of cells/ml, then with the average specific growth rate.

The binomial/non-linear interpolation or probit method was used to calculate the EC50 values. The no observed effect concentration (NOEC) is the highest concentration of the test substance that allowed at least 90% of control growth. This definition of the NOEC differs slightly from the TSCA test guidelines.

RESULTS

Exposure of the algae to the test substance resulted in the following growth after 96 hours:

Nominal concentrations (mg/l)	Average cells/ml	% of Control
0 (control)	1,945,000	-
0.3	1,708,000	88
3.3	86,000	4
33	<10,000	<1
330	<10,000	<1
3300	<10,000	<1

EC50 value with 95% confidence limits calculated using the number of cells/ml:

EC50 (72hr) 0.88 mg/l (0.3 – 3.3 mg/l)

EC50 (96hr) 0.83 mg/l (0.3 - 3.3 mg/l)

EC50 value with 95% confidence limits calculated using average specific growth rate:

EC50 (72hr) 2.7 mg/l (2.1 - 3.4 mg/l)

EC50 (96hr) 2.5 mg/l (2.0 - 3.2 mg/l)

NOEC (96hr) < 0.3 mg/l, calculated using the number of cells/ml

NOEC (96hr) 0.3 mg/l, calculated using average specific growth rate

Control response was satisfactory: the algal population grew well, resulting in an average of 1,945,000 cells/ml after 96 hours.

No biological effects (size differences, unusual cell shapes, colors, flocculations, adherence of cells to test containers or aggregation of cells) were noted during the test.

Remarks: The 3300 mg/l test concentrations were slightly cloudy at the start of the test and there was floating material in the 330 and 3300 mg/l solutions at 24, 48, 72 and 96 hours. No other insoluble material was noted during the test.

The pH of the test media was slightly affected by the test substance at all concentration above 3.3 mg/l.

The determination of whether toxic effects were algistatic or algicidal was performed at the end of the study. A 0.5 ml aliquot of test media from each 33 mg/l vessel was transferred to a flask containing 100 ml of fresh media and incubated under test conditions for 216 hours. Algae did not increase from initial density of less than 10,000 cells/ml, indicating that the effect of the test substance at 33 mg/l was algicidal rather than algistatic.

CONCLUSIONS

The 96-hour no observed effect concentration (NOEC) is less than 0.3 mg/l when determined using the number of cells/ml and 0.3 mg/l when determined using the average specific growth rate.

DATA QUALITY

Method follows OECD guideline.

Information taken from the test report.

GLP study.

REFERENCES

Acute Toxicity of the Water Accommodated Fraction (WAF) of OS # 43247K to the Fresh Water Algae *Selenastrum capricornutum*. – Range finding test

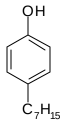
Report Number 1394-LU

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.21 mg/l

ChV (96hr) 0.12 mg/l

Remark: log Kow used 5.01 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.21 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

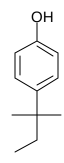
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

p- tert- Amylphenol

CAS No. 80-46-6



METHOD

Acute Toxicity to Non-Standard Species

Test type: unknown

GLP: No data

Year study performed: 1981

Species: Crangon septemspinosa (Shrimp)

Analytical monitoring: No data

Exposure period: 96 hours

RESULTS

LC50 (96hr) 0.3 mg/l

The LC50 (96 hour) was determined to be 0.3 mg/l.

CONCLUSIONS

The LC50 (96 hour) was determined to be 0.3 mg/l.

DATA QUALITY

Unknown

REFERENCES

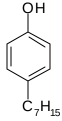
McLeese et al (1981) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.61 mg/l

ChV (21day) 0.043 mg/l

Remark: log Kow used 5.01 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.61 mg/l.

DATAQUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

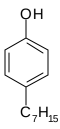
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Similar to that described in Section 1500.3 - Federal Hazardous Substances Act Regulations - 16 CFR

GLP: Data not available

Year study performed: 1982

Species: Sprague-Dawley strain rat.

Sex: Males and females.

Route of administration: Oral gavage.

Remarks: 5 male and 5 female albino rats of the outbred Sprague-Dawley strain weighing between 200 and 300 g each was employed in this study. The rats were deprived of food but not water overnight prior to dosing. Each animal was weighed and dosed by direct administration of the experimental material into the stomach by means of a syringe and dosing needle. The dose level administered was 0.2 g/kg, 2.0 g/kg and 5.0 g/kg. Following administration the animals were allowed food and water *ad libitum* for the 14-day observation period during which time the rats were observed for signs of toxicity and mortalities.

RESULTS

LD50: >0.2 g/kg and <2g/kg

0.2 g/kg – The animals were ruffled after 3 hours. They had dirty coats with urine stains and a bloody discharge around the nose and mouth within 24 hours. Between 12 and 24 hours, the animals were vocalising in high pitch squeaks. The dirty coats and discharge gradually improved and the animals appeared recovered by Day 3. They appeared normal throughout the remainder of the observation period. There were no remarkable gross pathologic findings.

2.0 g/kg - The animals appeared ruffled after 3 hours. Four animals died within 24 hours. The remaining animals were dirty, oily and depressed with a discharge around the facial orifices after 24 hours. These animals subsequently died on days 2 and 3. Gross pathology noted pale spleens and mottled liver.

5.0 g/kg – The animals were lethargic with diarrhoea and dirty, oily coats after 3 hours. Eight animals died within 24 hours. The remaining animals were moribund exhibiting a discharge around the facial orifices. They died within 48 hours. Gross pathologic examination revealed nothing remarkable.

CONCLUSIONS

LD50 = >0.2 g/kg and <2 g/kg.

Remarks:

Considering the relatively severe clinical signs that were seen at 0.2 mg/kg/day and consistent with those seen at higher dose which resulted in deaths, it would be unethical to do further testing to establish an actual LD50 value.

showing these signs died on Days 2 or 3.

At 0.2g/kg ruffled appearance was again seen after 3 hours, with dirty coats and blood discharge around the nose and mouth within 24 hours. Animals vocalised (high pitched squeaks) between 12 and 24 hours.

The OECD Guidance Document on the Recognition, Assessment, and Use of Clinical Signs as Humane Endpoints for Experimental Animals Used in Safety Evaluation (2000), Annex 3, states the following regarding failure to groom, piloerection and vocalisation:

Failure to groom—"The animal is definitely ill, and may be in severe pain and discomfort".

Piloerection – This can be a sign of not grooming and general ill health.

Vocalisation – If abnormal or persistent, may be indicative of a painful or distressful condition.

that clear toxicity was being seen at 0.2g/kg was the 21g group mean bodyweight loss seen for males between Day 1 and 7.

In view of the comments above it would be unethical to dose any further animals at or above 0.2g/kg in order to gain further information on the LD50.

DATA QUALITY

No data available on GLP.

Information taken from test report.

Purity of the test substance/decomposition: not known.

REFERENCES

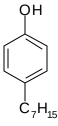
Biosearch Incorporated., P.O. Box 8598, Philadelphia, Pennsylvania 1910, USA. Acute Oral Toxicity in rats of the subject material. Project No. 82-3108A-1, 82-3108A-2 and 82-3108A.

13.2 ACUTE TOXICITY DERMAL

TEST SUBSTANCE

Heptyl derivs (*p*-heptylphenol)

CAS No. 72624-02-3



METHOD

Similar to those proposed in 40 CFR, Section 163.81-2, Federal Register, August 22, 1978 and subsequently modified in accordance with the revised EPA Pesticide Assessment Guidelines of Nov 1982.

GLP: Data not available

Year study performed: 1985

Species: New Zealand White rabbit.

Sex: Males and females.

Remarks: The test article was dosed as supplied, at a dose level of 2.0 g/kg. A group of 10 rabbits (5 male and 5 female) with healthy intact skin was used. Approximately 24 hours before testing, the fur was clipped from the backs of the test animals. All rabbits were weighed and the correct amount of test article was applied to approximately 10% of the body surface on each animal. The treated area was covered with a large porous gauze patch and wrapped with an impervious material to ensure that the animals did not ingest the test article. The dressings were removed after 24 hours and any excess material removed, where practical, using water or an appropriate solvent. The animals were observed for a 14-day period for signs of toxicity (systemic and topical) and for mortality.

RESULTS

LD50: >2.0 g/kg.

Males – Signs of necrosis and severe oedema were observed in 5/5 animals after unwrapping at 24 hours. Eschar was noted in 3/5 animals at 48 hours and in 2/5 hours at 72 hours. The eschar began to peel at 7 days in 3/5 animals, at 8 days in 1/5 animals and at 9 days in 1/5 animals. Other than the above skin observations, all animals appeared normal throughout the 14-day observation period. A loss of body weight was noted in 1/5 animals at 7 and 14 days. There were no gross abnormalities noted in the animals necropsied at the conclusion of the 14-day observation period.

Females – Signs of necrosis and severe oedema were observed in 5/5 animals after unwrapping at 24 hours. Eschar was noted in 5/5 animals by 48 hours, which began to peel in 4/5 animals at 8 days and in 1/5 animals at 10 days. Other than a loss of body weight at 7 days, no adverse symptoms preceded the death of one animal on day 12. Other than the above skin observations, the remaining 4/5 animals

appeared normal throughout the observation period. A loss of body weight was also noted in 2/5 animals at 7 days and in 1/5 of the remaining animals at 14 days. Diarrhoea, signs of dehydration and no formed faecal material in the lower gastrointestinal tract were noted in the animal found dead at 12 days. No gross abnormalities were noted in the animals necropsied at the conclusion at the 14-day observation period.

CONCLUSIONS

LD50 = > 2.0 g/kg.

DATA QUALITY

No data available on GLP.

Information taken from test report.

Purity of the test substance/decomposition: Not known.

REFERENCES

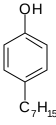
Biosearch Incorporated., P.O Box 8598, Philadelphia, Pennsylvania 1910, USA, Acute Dermal Toxicity, Single level - Rabbits. Project No. 85-4762A.

13.3 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Similar to that described in Section 1500.41, Federal Hazardous Substances Act Regulations - 16 CFR - P.124.

GLP: Data not available

Year study performed: 1982

Species: New Zealand White rabbit

Remarks: A group of six New Zealand White rabbits was clipped over a wide area. One side of the animals' backs was abraded at one site with a lancet sufficiently deep to penetrate the stratum corneum but not enter the derma to produce bleeding. The skin of the other side was allowed to remain intact. A 0.5 ml portion of material was applied to an abraded and an intact skin site on the same rabbit. Gauze patches were then placed over the treated areas and an impervious material was wrapped snugly around the trunks of the animals to hold the patches in place. The wrapping was removed at the end of the twenty-four hour period and the treated areas were examined. Readings were also made after 72 hours. The Draize method of scoring was employed.

RESULTS

Intact and abraded skin:

Erythema and eschar formation – scorings indicate severe erythema (beet redness) to slight eschar formation (injuries in depth) at the 24 and 72 hour observation periods.

Oedema formation – scorings indicate that slight oedema (edges of area well defined by definite raising) was noted (24 hour period) reducing to very slight oedema at the 72 hour period.

CONCLUSIONS

Based on the Draize method of scoring the test material is classified as a primary skin irritant.

DAT A QUALITY

No data available on GLP.

Information taken from test report.

Purity of the test substance/decomposition: Not known.

REFERENCES

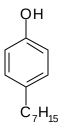
Biosearch Incorporated., P.O Box 8598, Philadelphia, Pennsylvania 1910, USA, Primary Skin Irritation - Rabbits. Project No. 82-3108A.

13.4 ACUTE TOXICITY EYE IRRITATION

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Similar to that described in Section 1500.42, Federal Hazardous Substances Act Regulations - CFR 16 P. 125.

GLP: Data not available

Year study performed: 1982

Species: Albino rabbit.

Remarks: Six healthy young adult albino rabbits were used in this study. 0.1 ml of the experimental material was instilled into the right eyes of the test animals while the other eyes remained untreated to serve as controls. The test material was not washed from the eyes. The treated eyes were examined at 1, 2, 3, 4, 7, 14, and 21 days following instillation of the test material into the eyes. Interpretation of the results was made in accordance with the Draize scale of scoring ocular lesions.

RESULTS

Using the Draize method of scoring the test material is classified as a primary ocular irritant.

CONCLUSIONS

The test material is classified as a primary ocular irritant.

DATA QUALITY

No data available on GLP.

Information taken from test report.

Purity of the test substance/decomposition: Not known.

REFERENCES

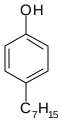
Biosearch Incorporated., Primary Eye Irritation - Rabbits. Project No. 82-3108A.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY)

TEST SUBSTANCE

Heptyl derivs (p-heptylphenol)

CAS No. 72624-02-3



METHOD

Ames, et al., 1975; Bridges, 1972; Green and Muriel, 1976; Maron and Ames, 1983; Maron, et al., 1981

GLP: (N)

Year study performed: 1993

Species/strain: Salmonella typhimurium TA 1535, TA 1537, TA 1538, TA 98 and TA 100 and Escherichia coli WP2uvrA.

Metabolic activation: S-9 mix (with and without metabolic activation) 6% Aroclor 1254 – induced male rat liver and homogenate.

Test 1. Salmonella (5 strains) 0.05, 0.167, 0.5, 1.67, 5.0, 16.7 µg/plate (+/- S9)

E.coli 0.167, 0.5, 1.67, 5.0, 16.7, 50.0 μg/plate

Test 2. Repeat TA 1538 + S9 mix

Repeat 6 strains – S9 mix

TA 1535, 1537, 98 and 100: 1.67, 5.0, 16.7, 50.0, 167, 500 μg/plate with S9 mix

Test 3. TA 1535, 1537, 98 and 100 (+S9)

0.5, 1.67, 5.0, 16.7, 50 and 100 µg/plate

WP2uvrA: 0.167, 0.5, 1.67, 5.0, 16.7, 50.0 and 100 µg/plate (+S9)

RESULTS

produced inhibited growth or complete toxicity in strains TA 1538 and TA 100 at doses ≥ 5.00 µg/plate and in strain WP2uvrA at doses ≥ 16.7 µg/plate.

Based upon these findings, the test substance was evaluated in triplicate cultures in all five *Salmonella* strains at doses of 0.0500, 0.167, 0.500, 1.67, 5.00 and 16.7 μ g/plate and in strain WP2 *uvrA* at doses of 0.167, 0.500, 1.67, 5.00, 16.7 and 50.0 μ g/plate in the presence and absence of S-9 mix. Six dose levels of the test substance were evaluated with and without S-9 mix in the event of unacceptable toxicity at the highest dose levels evaluated in the mutation assay.

The test substance was found to be freely soluble at all doses evaluated. Inhibited growth was observed in all tester strains at doses of 0.500, 1.67, 5.00, 16.7 and/or 50.0 μg/plate with S-9 mix, and in strain TA 1538 at doses of 5.00 and 16.7 μg/plate without the S-9 mix. Revertant frequencies for all doses of the test substance in all tester strains with and without S-9 mix approximated or were less than those observed in the concurrent negative controls. The test substance was re-evaluated in the confirmatory assay under identical conditions in strain TA 1538 with S-9 mix and in all six tester strains without S-9 mix. Based upon the limited toxicity observed in the previous trial, however, the test substance was re-evaluated in strains TA 1535, TA 1537, TA 98 and TA100 and WP2uvrA at doses of 1.67, 5.00, 16.7, 50.0, 167 and 500 μg/plate with S-9 mix. The test substance was again found to be freely soluble at all doses evaluated, and inhibited growth was observed in all tester strains at the highest two or three doses evaluated with and without S-9 mix. Revertant frequencies for all doses of the test substance in all five *Salmonella* tester strains with S-9 mix and in all six tester strains without S-9 mix approximated or were less than control values. A statistically significant increase in revertant frequency, to approximately 2.6-fold control values, was observed in tester strain WP2uvrA at a dose of 1.67 μg/plate. However, this increase apparently was not dose dependent.

Therefore, the test substance was re-evaluated in a third trial in strains TA 1535, TA 1537, TA 98 and TA 100 at doses of 0.500, 1.67, 5.00, 16.7, 50.0, and 100 μ g/plate and in strain WPuvrA at doses of 0.167, 0.500, 1.67, 5.00, 16.7, 50.0 and 100 μ g/plate with S-9 mix. The test substance again was found to be freely soluble at all doses evaluated and inhibited growth was observed in all tester strains at doses of 16.7, 50.0 and/or 100 μ g/plate with S-9 mix. Revertant frequencies for all doses of the test substance in tester strains TA 1535, TA 98, TA 100 and WP2uvrA approximated or were less than control values. A statistically significant increase in revertant frequency, to approximately 2.1-fold control values, was observed in tester strain TA 1537 at a dose of 1.67 μ g/plate. However, this increase apparently was not dose dependent. Thus the slight increases observed in strains TA1537 and WP2uvrA in the confirmatory assay or final retest are considered to be statistical aberrations due to random fluctuation of the spontaneous revertant frequencies.

All positive and negative control values in all assays were within acceptable limits.

All strains were treated in the presence and absence of metabolic activation with the test substance dissolved in dimethyl subhoxide (DMSO). Triplicate cultures of each strain were evaluated with the appropriate solvent in the presence and absence of S9 to serve as negative (solvent) controls. Triplicate cultures of each strain were also evaluated with known positive control chemicals. Positive control evaluated in the absence of S-0 mix were specific for each strain and included: TA 1535 and TA 100-sodium azide (10.0 μ g/plate); TA 1537- 9-aminoacridine (150 μ g/plate); TA 1538 and TA 98- 2-nitrofluorene (5.00 μ g/plate); and WP2uvrA- N-Ethyl-N-nitrosoguanidine

(ENNG) (2.00 μ g/plate). 2-Anthramine (2.00 μ g/plate) was evaluated in all five *Salmonella* tester strains, and in strain WP2uvrA (80.0 μ g/plate) in the presence of S-9 mix.

CONCLUSIONS

The test substance was considered to be non-mutagenic in this test system.

Not a GLP study.

Information taken from test report.

Purity of the test substance/decomposition: not known.

REFERENCES

Pharmakon USA., P.O Box 609, Waverley, Pennsylvania 18471-0609, USA, PA, Ames/Salmonella-E.coli liquid pre-incubation assay on OS #43247H. PH 301-LU-023-93.

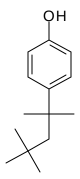
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Melting point: 79 - 82°C.

CONCLUSIONS

The test substance has a melting point range of 79 - 82°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

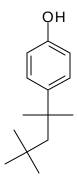
REFERENCES

Occupational Health Service Inc., New York (USA), Rev. 21.02.91 (CD-ROM). as cited in IUCLID database.

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Boiling point: 280 - 283°C at 101300 Pa (760 mm Hg).

CONCLUSIONS

The test substance has a boiling point range of 280 - 283°C at 101300 Pa.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

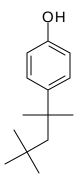
REFERENCES

Sax, Dangerous properties of Ind. Materials, 7th edition, 1989 as cited in IUCLID database.

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1989.

RESULTS

Vapor pressure: 0.21 Pa @ 20℃.

CONCLUSIONS

The test substance has a vapor pressure of 0.21 Pa @ 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

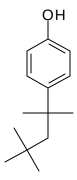
Huels-Produktdatenblatt "Octylphenol PT", Artikel-Nr. 002325, 1989 as cited in IUCLID database.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



METHOD

Method/guideline followed: OECD Guideline 107 "Partition Coefficient (n-octanol/water), Flask-shaking Method".

GLP (Y/N): no.

Year study performed: 1993.

RESULTS

Log Kow: 4.12 at 20.5°C.

CONCLUSIONS

The test substance has a log Kow of 4.12 at 20.5°C.

DATA QUALITY

Method follows OECD guideline. Temperature but not test substance purity given.

Information taken from IUCLID database.

REFERENCES

Ahel, M. and Giger, W.: Partitioning of alkylphenols and alkylphenol polyethoxylates between water and organic solvents. Chemosphere 26, 1471 – 1478 (1993) as cited in IUCLID Database.

OTHER

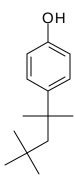
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



METHOD

Method/guideline followed: generator column.

GLP (Y/N): yes.

Year study performed: 1984.

RESULTS

Water solubility: 17 - 19 mg/l at 22°C.

CONCLUSIONS

The solubility of the test substance in water is 17 - 19 mg/l at 22°C.

DATA QUALITY

GLP study.

Information taken from IUCLID database.

REFERENCES

Analytical Bio-Chemistry Laboratories, Inc.:7200 ABC Lane, P.O Box 1097, Columbia, Missouri 65205: Method validation and solubility of octylphenol in aquatic test waters, unpublished test report No. 31914, December 1984 as cited in IUCLID Database.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: 42.4×10^{-12} cm³/molecule-sec.

Half-life: 3.03 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 3.03 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

ADDITION

A half-life of 14 days (in water) was measured by Waern (2000). Method and results not available.

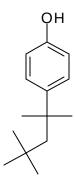
Waern (2000) and SIDS (2000) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for The UK Environment Agency by WRc-NSF.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

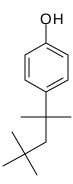
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:206Data temperature (°C):25Water solubility (mg/l):18Vapor pressure (Pa):0.21Log Kow:4.12Melting point (°C):80.5

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	3.61
Soil	86.9
Water	7.45
suspended sediment	0.060
fish	0.0049
Sediment	1.93

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

OTHER - ADDITION

Using Mackey Level III Type Fugacity model, the following was calculated:

Release 100% to water:

In Air: 1.2%

In Water: 77.9%

In Sediment: 17.8%

In Soil: 3.1%

SIDS (2000) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for The UK Environment Agency by WRc-NSF.

9.1 BIODEGRADATION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9

METHOD

METHOD

Method/guideline followed: OECD Guideline 302 C "Inherent Biodegradability: Modified MITI Test (II)"

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1981.

Contact time: 28 days.

Inoculum: Mixed population of non-adapted organisms from activated sewage sludge.

Concentration: 30 mg/l (equivalent to 100 mg/l dry weight)

RESULTS

Percentage degradation: 0% after 28 days.

CONCLUSIONS

p-tert-Octylphenol was not inherently biodegradable under the conditions of the study.

DATA QUALITY

Not a GLP study.

Purity of test substance not specified.

Recognised study design used.

REFERENCES

Safepharm Laboratories Ltd., P.O Box 45, Derby, DE1 2BT, U.K. Assessment of inherent biodegradability: Modified MITI (II), unpublished report No. S0052/E584, November 1991 as cited in IUCLID database.

SIDS 1994 and IUCLID 2000

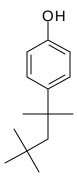
9.2 BIODEGRADATION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9

Purity: 95%



METHOD

Method/guideline followed: BOD insoluble substances Test (ISO 10708, in preparation)

Test type: aerobic.

GLP (Y/N): yes.

Year study performed: 1991.

Contact time: 28 days.

Inoculum: non-adapted activated sewage sludge.

Concentration: 27.5 mg/l (related to test substance).

RESULTS

Percentage degradation: 20% after 28 days.

CONCLUSIONS

*p-tert-*Octylphenol was not readily biodegradable under the conditions of the study. This material is considered to be inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

GLP study.

REFERENCES

Huels-Bericht Nr. BO-91/5, 1991 (unveroeffentlicht) as cited in IUCLID database.

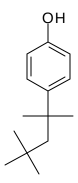
9.3 BIODEGRADATION - ADDITION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9

Purity: 99.64%



METHOD

Method/guideline followed: OECD Guideline 301B

Test type: aerobic.

GLP (Y/N): Yes

Year study performed: 1999.

Contact time: 35 days.

Inoculum: Domestic activated sludge from a sewage treatment plant where a high concentration of nonylphenol ethoxylate (> $600 \mu g/l$) was found.

Concentration: 30 mg/l (equivalent to 100 mg/l dry weight)

RESULTS

A mean net carbon dioxide value of 69.9% of the theoretical amount (based on titration measurements) was measured following 35 days exposure.

CONCLUSIONS

69.9% of theoretical CO₂ produced following 35 days.

While, based on these results, 4 tert-octylphenol could not be characterised as "readily biodegradable" (under which OECD definition requires 60% or greater carbon dioxide with 28 days, with this value being achieved within a 10-day window), it was extensively mineralised (> 60 % CO₂) during the 35 day study. In addition, carbon dioxide and dissolved organic carbon concentration confirmed extensive removal (>90%) of the initial test substance organic carbon by mineralisation, adsorption and biomass incorporation. The results from this study suggest that micro-organisms may need a period of adaptation before degradation of 4-tert-octylphenol.

DATA QUALITY

OECD and GLP compliant.

Summarised in Targeted Risk Assessment Report.

REFERENCES

Gledhill (1999) as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency.

OTHER

Johnson et al (2000) investigated the potential for 4-tert-octylphenol to biodegrade in some English rivers. Using laboratory microcosms, half-lives of 7 to 50 days were obtained for water samples. Shorter half-lives were generally seen in more urban/ industrialised rather than upland/rural areas. However, even then, half-lives varied within the river for samples of river water taken at different times, although a similar degradation rate was noted for a range of concentrations from 0.3 to $100~\mu g/l$. Hence this work demonstrated that 4-tert-octylphenol could be degraded in river samples taken from a range of urban and rural reaches (with the possible exception being a sample taken from an upland stream). No degradation was observed over 83 days when bed sediments were spiked with 4-tert-octylphenol and incubated under anaerobic conditions.

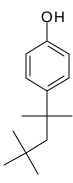
Johnson et al. (2000), as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency.

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Purity of the test substance stated as 99.34%.

METHOD

American Public Health Association. 1980. Standard Methods for the Examination of Water and Wastewater. 15th ed. Washington, DC. 1134 p.

Committee on Methods for Toxicity Tests with Aquatic Organisms (C. E. Stephan, Chairman). 1975. Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and Amphibians. Environmental Protection Agency, Ecological Research Series EPA-660/3-75-009, April, 1975. 61 p.

Test type: 96-hour flow-through toxicity test

GLP: (Y)

Year study performed: 1984

Species: fathead minnow (Pimephales promelas)

Analytical procedures: The concentrations of octylphenol in water were determined by extraction and HPLC methods on days 0 and 4 of the study.

Test substance preparation: A proportional diluter system was used for the intermittent introduction of octylphenol and diluent water into the test aquarium. Parameters of temperature, dissolved oxygen and pH were measured at 0, 48 and 96 hours in the control, low and high test concentrations with remaining fish at the time of sampling:

Temperature $15 - 20^{\circ}$ C

Dissolved oxygen 9.2 – 10.2 ppb

pH 7.8 - 8.3

Hardness (CaCO₃) 225 - 275 ppm

A diluter stock of 14,500 mg/l in acetone was prepared of the test compound for the definitive study.

The test was initiated by stratified random assignment of 20 fathead minnow to each test aquarium after the test solutions had been flowing through the aquarium for at least 24 hours. A control received 0.1 ml of acetone that was equivalent to that received in the highest test concentration.

Exposure period: 96 hours

Statistical methods: Binomial Method

Moving Average Method

Probit Method

RESULTS

Nominal concentrations: 0.047, 0.091, 0.18, 0.39 and 0.70 mg/l.

Mean measured concentrations: 0.041, 0077, 0.15, 0.34 and 0.63 mg/l. These values ranged from 83 to 90% of nominal concentration.

LC50 values with 95% confidence limits and no-observed effect concentration, based on mean measured concentrations:

LC50 (24hr) 0.29 mg/l (0.15 - 0.63 mg/l)

LC50 (48hr) 0.25 mg/l (0.15 - 0.34 mg/l)

LC50 (72hr) 0.25 mg/l (0.15 - 0.34 mg/l)

LC50 (96hr) 0.25 mg/l (0.15 - 0.34 mg/l)

NOEC (24hr) 0.15 mg/l

NOEC (48hr) 0.077 mg/l

NOEC (72hr) 0.077 mg/l

NOEC (96hr) 0.077 mg/l

The fish were observed for mortality and abnormal effects daily for the 96-hour test period. Abnormal effects monitored included surfacing, loss of equilibrium, color changes, erratic swimming and lying on the bottom of the chamber. Mortality, surfacing, loss of equilibrium, dark discoloration and/or quiescence were observed in level 3, 4 and 5 during the 96 hour study.

re

	24	48	72	96
Solvent control	0	0	0	0
(Level 1) 0.047	0	0	0	0
(Level 2) 0.091	0	0	0	0
(Level 3) 0.18	0	0	0	0
(Level 4) 0.39	65	90	90	90
(Level 5) 0.70	100	100	100	100

CONCLUSIONS

Based on the lack of mortality and abnormal effects, the results indicated a 96-hour LC50 and 95% confidence interval of 0.25 mg/l (0.15 - 0.34 mg/l) and a no-observed effects concentration of 0.077 mg/l for octylphenol.

Remarks: The primary objective of this study was to define the acute toxicity of octylphenol by determining the LC50's during the course of the test. A second objective of the study was to provide a lethal threshold (incipient LC50) for either fathead minnow (*Pimephales promelas*) or rainbow trout (*Salmo gairdneri*) depending on which was the single most sensitive species. It was concluded that the rainbow trout appeared to be the more sensitive species; therefore the fathead minnow study was terminated after 96 hours (see below). However, a lethal threshold appeared to be reached after 48 hours and was estimated to be 0.25 mg/l.

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Dynamic 96-Hour Acute Toxicity of Octylphenol to Fathead Minnows (*Pimephales promelas*). Final Report # 31910

OTHER

Results of the dynamic 14-day rainbow trout (*Oncorhynchus mykiss*) study- LC50s with their 95% confidence interval and no observed effect concentrations:

LC50 (1day) 0.45 mg/l (0.32– 0.71 mg/l)	NOEC(1day) 0.17 mg/l
LC50 (2days) 0.37 mg/l (0.17-0.71 mg/l)	NOEC(2day) 0.17 mg/l
LC50 (3days) 0.24 mg/l (0.17-0.32 mg/l)	NOEC(3day) 0.17 mg/l
LC50 (4days) 0.21 mg/l (0.17 –0.32 mg/l)	NOEC(4day) 0.17 mg/l
LC50 (5days) 0.20 mg/l (0.084 -0.32 mg/l)	NOEC(5day) 0.17 mg/l
LC50 (6days) 0.17 mg/l (0.084 -0.32 mg/l)	NOEC(6day) 0.17 mg/l
LC50 (7-8days) 0.14 mg/l (0.084 –0.17 mg/l)	NOEC(7-8day) 0.17 mg/l
LC50 (9days) 0.13 mg/l (0.084 –0.17 mg/l)	NOEC(9day) 0.17 mg/l
LC50 (10-14day) 0.12 mg/l (0.084 -0.17 mg/l)	NOEC(10-14day) 0.17 mg/l

OTHER - ADDITION

A similar 96 hour LC50 value of 0.26 mg/l was measured for the golden orfe (Leuciscus idus) in a semi-static test, in which concentrations were measured and the study was carried out to GLP guidelines (IUCLID 2000). In the same study an LC0 of 0.21 mg/l and an LC100 of 0.390 mg/l were also obtained. A higher 48-hour LC50 was found for the same species in another study based on nominal concentrations and not carried out to GLP. (IUCLID, 2000).

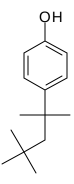
IUCLID, (2000), as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

10.2 EARLY LIFE STAGE TEST

TEST SUBSTANCE

*p- tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Purity of the test substance given as 99.22%

METHOD

Proposed Recommended Bioassay Procedure for Egg and Fry Stages of Fresh Water Fish. ASTM Standard Practice for Conducting Toxicity Tests on the Early Life Stages on Fish.

Test type: 60 day flow -through post-hatch early life stage toxicity test

GLP: (Y)

Year study performed: 1986

Species: Rainbow trout Salmo gairdneri

Analytical procedures: The actual concentration of octylphenol was measured by extraction and HPLC method.

Test substance preparation: A two-liter proportional diluter system was used for the intermittent introduction of an acetone solution of octylphenol to four replicate test chambers per concentration. Flow - splitting cells divided each of the five test concentrations, acetone solvent control and control solutions into the respective replicate test chambers. Aerated well water was delivered to the glass aquarium at an average rate of approximately 50 ml/minute per replicate.

Water quality parameters were measured at day 0, 1 and on every \mathcal{T}^h day during the study:

Temperature: 15-20 °C

Dissolved oxygen: 9.21 – 10.1 ppm

pH: 7.8 - 8.3

Hardness (CaCO₃): 225 - 275 ppb

Alkalinity (CaCO₃): 325 - 375 ppb

Exposure period: 60 days

Statistical methods: The statistical data were analysed by Systat® computer program (Version 2.1). Growth data, using the individual per replicate, were analysed using two-way analysis of variance with an interaction model. One-way analysis of variance (ANOVA) calculations were used to determine whether significant difference existed between the controls and treatment levels, followed by Tukey's HSD multiple means comparison test.

RESULTS

Nominal concentrations: 0.0062, 0.012, 0.025, 0.050 and 0.10 mg/l.

Mean measured concentrations: $0.0061 (\pm 0.0024)$, $0.011 (\pm 0.0025)$, $0.022 (\pm 0.0046)$, $0.051 (\pm 0.0078)$ and $0.091 (\pm 0.0079)$ mg/l. Values ranged from 96 to 125% of nominal concentrations.

MATC (Maximum Acceptable Toxicant Concentration): 0.0061 – 0.011 mg/l

NOEL 0.0061 mg/l

Morphological and behavioural abnormalities of quiescence, curvature of the spine and loss of equilibrium occurred consistently in the highest three test concentrations prior to mortality. Surfacing and erratic swimming were occasionally noted before mortality occurred.

Hatch-ability of rainbow trout eggs after 20 days of continuous exposure to the test substance was not significantly affected when compared to control. Survival of fry between hatch and 60 days of exposure to the three highest test concentrations was significantly reduced when compared to control.

Rainbow trout fry growth was significantly reduced after 35 and 60 days of exposure to the four highest test concentrations. Growth of trout fry, as measured by wet weight after 60 days of exposure, was significantly reduced in the three highest test concentrations.

CONCLUSIONS

The maximum acceptable toxicant concentration of the test substance was found to be greater than 0.0061 mg/l and lower than 0.011 mg/l.

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Early Life Stage Toxicity of *para-tert*-octylphenol to Rainbow Trout (*Salmo gairdneri*) in a Flow-Through System.

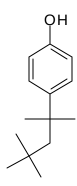
Final Report #34452, Analytical Bio-Chemistry Laboratories, Inc, 7200 ABC Lane, P.O Box 1097, Columbia, Missouri 65202, USA.

10.3 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed:2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.29 mg/l

ChV(30 day) 0.041 mg/l

ChV (90 day) 0.008 mg/l

Remark: log Kow used 5.28 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.29 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

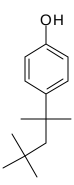
Calculation performed for this HPV submission.

10.4 CHRONIC TOXICITY TO FISH – SUMMARY ADDITION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

SUMMARIES OF THREE ADDITIONAL STUDIES

Embryotoxicity tests were carried out by Gray and Metcalfe (1999) on the Japanese medaka (*Oryzias latipes*). Fish were exposed from day 0 (fertilisation) to day 17 (swim-up) using a static non-renewal procedure. However LC50 values (following 17 days exposure) for the three replicates had a significantly wide range with calculated values of 450, 830, and 94 µg/l (based on nominal concentrations) reported. Developmental abnormalities observed in embryos and larvae ranged from circulatory problems to failure to inflate swim bladders.

Ashfield et al. (1995) investigated the effect of exposure to octylphenol (not stated if 4-tert-octylphenol) on the growth rate of rainbow trout ($Oncorhynchus\ mykiss$). Octylphenol was found to cause significantly increased growth of juvenile rainbow trout at 1 µg/l, but at 10 and 30 µg/l, caused growth suppression when exposed for 35 days. These effects were still apparent one year after exposure was terminated. However, the paper from which this information was obtained gives no information on methods used, and therefore these values should be treated with caution.

A 60-day post-hatch early life stage toxicity study with rainbow trout has also been carried out under flow-through and GLP protocol conditions. Test concentrations were 6.1, 11, 22, 51 and 91 μ g/l based on measured concentrations and a NOEC of 6.1 μ g/l and an LOEC of 11 μ g/l were identified, although the toxic endpoint was not specified. (IUCLID, 2000).

REFERENCES

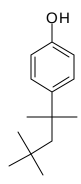
Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

The sample of the test substance was high purity grade.

METHOD

Based on:

Miller, W. E., J. C. Greene and T. Shiroyama. July 1978. The *Selenastrum capricornutum* Printz Algal Assay Bottle Test. Experimental Design, Application and Data Interpretation Protocol. Special Studies Branch, Corvallis Environmental Research laboratory, Corvallis, Oregon 97330. EPA-600/9-78-018

OTS Algal Acute Toxicity Test, August 1982. EG-8, ES-5

ASTM Proposed Standard practice for conducting short-term toxicity tests with fresh water and salt water algae. E-47.01, Draft 5, March 1983

Test type: 96-hour static algal assay

GLP: (Y)

Year study performed: 1984

Species: Green algae - Selenastrum capricornutum Printz

Exposure period: 96 hours, from 6th to 10th September

Test substance preparation: An initial definitive stock solution of octylphenol was made by weighing 100 mg into a 10 ml volumetric flask and diluting with triethylene glycol. Appropriate additions were made of the stock solution with triethylene glycol so that 0.1 ml total triethylene glycol was added to each test flask, which gave the appropriate test concentrations.

Analytical procedures:

Statistical methods:

Stephan, C. E., K. A. Busch, R. Smith, J. Burke and R. W. Andrews. 1978. A computer program for calculating an LC50. U.S. Environmental Protection Agency, Duluth, Minnesota.

One-way analysis of variance (ANOVA).

Multiple means comparison test (Fischer's LSD).

The test vessels were incubated for 96 hours at 24-25% under "cool-white" fluorescent light and constant shaking. Light intensity was maintained at $400\pm10\%$ ft-c and shaker speed was 100 rpm. Temperature and light readings measured throughout the test were within acceptable limits. Log phase growth was confirmed at 96 hours with a count of 9 x 10^5 cells/ml in the control. A 96-hour range-finding study was conducted to determine the concentration range in the definitive study. The flasks were prepared in triplicate for each test concentration, solvent control and control. The dilution water was sterile enriched media adjusted to a pH of 7.5 ± 0.1 .

RESULTS

Nominal concentrations: 1.0, 1.8, 3.2, 5.6 and 10 mg/l

EC50 value with 95% confidence limits and no-effect level:

EC50 (96hr) 1.9 mg/l (1.0 - 2.7 mg/l)

NOEC (96hr) < 1.0 mg/l

CONCLUSIONS

The one-way ANOVA results indicated a 96-hour, no observed effect concentration of less than 1.0 mg/l. All test concentrations were significantly different from the control (α = 0.05).

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Acute Toxicity of Octylphenol to *Selenastrum capricornutum* Printz. Analytical Bio-Chemistry Laboratories, Inc, 7200 ABC Lane, P.O Box 1097, Columbia, Missouri 65202, USA.

Static acute bioassay report # 31913

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE) - ADDITION

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9

IUCLID Identification Number: 140-66-9

METHOD

Test type: 72-hour static algal assay

GLP: (unknown)

Year study performed: unknown

Species: Green algae – Scenedesmus subspicatus

Exposure period: 72 hours

RESULTS

Based on effects on growth rate:

EC10 (72hr) 0.3 mg/l

EC50 (72hr) 1.1 mg/l

EC90 (72hr) 4.2 mg/l

CONCLUSIONS

The results indicated a 72-hour, EC50 of 1.1 mg/l.

DATA QUALITY

IUCLID 2000

REFERENCES

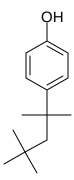
IUCLID (2000) as cited in Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

11.3 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.13 mg/l

ChV (96hr) 0.085 mg/l

Remarks: log Kow used 5.28 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.13 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

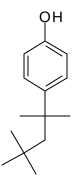
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Purity of the test substance stated as 99.34%.

METHOD

American Public Health Association. 1975. Standard Methods for the Examination of Water and Wastewater. 14th ed. Washington, DC. 1193 p.

Committee on Methods for Toxicity Tests with Aquatic Organisms (C. E. Stephan, Chairman). 1975. Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and Amphibians. Environmental Protection Agency, Ecological Research Series EPA-660/3-75-009, April, 1975. 61 p.

Test type: 48-hour flow-through toxicity test

GLP: (Y)

Year study performed: 1984

Species: Daphnia magna

Analytical procedures: The concentrations of octylphenol in water were determined by extraction and HPLC on days 0 and 2 of the study.

Test substance preparation: A proportional diluter system was used for the intermittent introduction of octylphenol and diluent water into the test chambers. The system contained seven sets of four replicate one-litre test chambers designated as control, solvent control and level 1 through level 5. Fbw-splitting chambers were utilised to thoroughly mix and divide each octylphenol concentrations for delivery to the test chambers. Water quality parameters of temperature, dissolved oxygen and pH were measured at 0 and 48 hours in the control, low and high test concentrations with remaining daphnids:

Temperature

 $15-20^{\circ}C$

Dissolved oxygen 9.2 – 10.2 ppb

pH 7.8 - 8.3

Hardness (CaCO₃) 225 - 275 ppm

A diluter stock of 24800 mg/l in acetone was prepared of the test compound for the definitive study.

Statistical methods: Binomial Method

Moving Average Method

Probit Method

RESULTS

Nominal concentrations: 0.072, 0.12, 0.25, 0.43 and 1.0 mg/l.

Mean measured concentrations: 0.063, 0.11, 0.19, 0.32 and 0.94 mg/l. These values ranged from 74 to 94% of nominal concentrations.

LC50 values with 95% confidence limits based on mean measured concentrations:

LC50 (24hr) 0.26 mg/l (0.19 - 0.32 mg/l)

LC50 (48hr) 0.27 mg/l (0.19 - 0.32 mg/l)

NOEC (48hr) 0.11 mg/l

The daphnids were observed for mortality and abnormal effects daily for the 48-hour test period. Abnormal effects monitored included surfacing, quiescence, loss of equilibrium, clumping of the daphnids together and lying on the bottom of the test chamber. Mortality, quiescence and/or daphnids on the bottom of the test chamber were observed in the 0.19, 0.32 and 0.94 mg/l test concentrations after 48 hours. The difference in the 24 and 48 hour survival of the 0.32 mg/l test concentration (90 and 82% respectively) was due to the recovery of 3 daphnids that appeared dead after 24 hours.

CONCLUSION

Based on the lack of mortality and abnormal effects, the results indicated a 48-hour no- observed effect concentration of 0.11 mg/l for octylphenol.

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Dynamic 48-Hour Acute Toxicity of Octylphenol to Daphnia magna. Final Report # 31912

OTHER - ADDITION

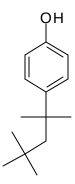
In two other studies using the same species, a 24-hour EC50 of 170 ug/l and a 48 hour lC50 of 90 ug/l have been reported (IUCLID (2000) and Zou and Fingerman (1997) respectively). However, the first was not carried out to GLP and had no analytical monitoring, while no information on study conditions was provided in the second paper. Both studies should therefore be used with care. As cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SHRIMP) - ADDITION

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Purity of the test substance: unknown

METHOD

Test type: 96-hour semi-static toxicity test

GLP: (unknown)

Year study performed: 1998

Species: Gammerus pulex (freshwater shrimp)

Analytical procedures: Analytical confirmation of test solutions.

RESULTS

First and second instar nymphs of the freshwater shrimp exhibited a 96-hour EC50, based on immobilisation of 0.013 mg/l and a corresponding LC50 of 0.0196 mg/l. There was also a clear time-dependence of toxicity although toxicity appeared to begin to plateau by 96 hours.

EC50 (96 hour) 0.013 mg/l

LC50 (96 hour) 0.019 mg/l

These results were based on immobilisation.

CONCLUSION

These freshwater shrimp had a 96 hour EC50 of 0.013 mg/l and a 96 hour LC50 of 0.019 mg/l.

DATA QUALITY

Deemed to be appropriate/ good quality (although not to standard guidelines) by UK RAR Process.

REFERENCES

Sims and Whitehouse (1998) as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency, as well as "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

OTHER - ADDITION

For saltwater species, a series of acute toxicity data for the opossum shrimp (*Americamysis bahia*) were located, providing 96-hour LC50 values ranging from 0.0534-0.1131 mg/l. (Cripe et al. (1989)) However, these data were taken from a study investigating the effects of different feeding regimes on the acute toxicity of 4 substances to the opossum shrimp, and were not carried out to standard guidelines. Thus, there were a number of variables which could influence the results, such as nutritional deficiency (direct physiological effect on susceptibility to toxicants) or surplus food in the test environment (possible effects on DO in a static test, again inducing physiological stress). While the procedures themselves appear to have been adequately carried out for the endpoints desired, and well reported, there were obvious difficulties in using these data for direct comparison with other standard invertebrate toxicity data, such as lethality data obtained for the freshwater amphipod *gammerus pulex*. However, it was noted that the range of LC50 values observed were broadly similar to those of the freshwater amphipod.

Cripe et al (1989) as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

The other data located for a saltwater species were for the fiddler crab (*Uca pugilator*) where after 3 and 7 days exposure to 4-tert-octylphenol at a concentration of 10 mg/l, enzymatic changes were recorded (Zou and Fingerman, 1999). The changes recorded in this study may potentially be due to endocrine disruption induced by octylphenol. The concentrations inducing these effects were significantly higher (at least an order of magnitude) than those causing effects on more traditional endpoints in other aquatic invertebrates. (Changes in levels of enzyme chitobiase, which is involved in the breakdown of chitinous cuticle during moulting, following stimulation by the ecdysteroid moulting hormones.) These results are useful as there are few data for studies on the ecdysteroid system in aquatic organisms and the consequences of xenobiotic-induced disruption. However, they cannot be taken at this stage, as evidence of true endocrine disruption in the crab resulting from PTOP exposure.

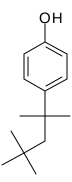
Zou and Fingerman, 1999 as cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency

12.3 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Purity of the test substance given as 99.34%.

METHOD

American Society for Testing and Materials. 1981. Proposed Standard Practice for Conducting *Daphnia magna* Chronic Toxicity Tests in Flow-Through Systems. Draft No. 3, 1981.

Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and Amphibians. EPA, Ecological Research Series EPA-660/3-75-009, 1975. 61p.

U. S. EPA 1976. TSCA. Public Law 94-469. Federal Register, 1976. 2003 – 20521

Test type: Flow though toxicity test

GLP: (Y)

Year study performed: 1988

Species: Daphnia magna

Exposure period: 21 day

Analytical procedures: The concentrations of the test substance in water were determined by HPLC on days 0, 4, 7, 14 and 21.

Test substance preparation: A proportional diluter system was used for the intermittent introduction of octylphenol and diluent water into the test chambers. The system contained seven sets of four replicate (total of 40 daphnids) one-litre test chambers designated as control, solvent control, and level 1 through level 5. Flow-splitting chambers were utilised to thoroughly mix and divide each octylphenol concentration for delivery to the test chambers. Aerated water was delivered to each chamber at an average rate of 4.1 ml/min.

Water quality parameters of dissolved oxygen and pH were measured at day 0, 4, 7, 14 and 21.

Temperature: 20°C

Dissolved oxygen: 7.5 - 9 ppb

pH: 7.8 - 8.3

TOC and suspended solids (routinely analysed on a monthly basis):

Hardness (CaCO₃): 206 - 275 ppm

Alkalinity (CaCO₃): 224 - 336 ppm

TOC: 1.4 ppm (0.9 - 2.8 ppm)

Suspended solids: 0- 1.3 ppm

All stock solutions were corrected for compound purity and prepared in acetone.

Statistical methods:

One-way analysis of variance (ANOVA)

Multiple means comparison test

Dunnett C. W. 1985. A Multiple Comparison Procedure for Comparing Several Treatments with a Control. J. Amer. Stat. Assoc. 50: 1096-1121

Stephan C. E., K. A. Busch, R. Smith, J. Burke and R. W. Andrews. 1978. A computer Program for Calculating an LC50. U.S. EPA, Duluth, Minnesota, pre-publication manuscript 1978.

Stephan, C. 1977. Methods for calculating an LC50, p. 65-84. In F. L. Mayer and J. L. Hamelink. Aquatic Toxicology and Hazard Evaluation. ASTM Special Technical Publication 643. ASTM Philadelphia.

RESULTS

Nominal concentrations: 0.030, 0.060, 0.12, 0.25 and 0.50 mg/l

Mean measured concentrations: 0.037, 0.062, 0.12, 0.23 and 0.51 mg/l. These values ranged from 92 to 123% of nominal concentrations.

EC50 (21day) 0.34 mg/l

NOEC (21 day) 0.037 mg/l (based on growth/survival)

MATC >0.037 and< 0.062 mg/l

Statistical analysis of survival for *Daphnia magna* after a 21-day exposure to octylphenol indicated that daphnid survival was significantly different ($P \le 0.05$) from the pooled controls in the mean measured concentration of 0.51 mg/l. All daphnids had died in the 0.51 mg/l test level by day 9 of the study. Therefore, no reproduction and length data were available for further statistical analysis. A day 21 EC50 was calculated to be 0.34 mg/l.

The mean young/adult/ reproduction day after 21 days were significantly affected in the mean measured exposure levels of 0.12 and 0.23 mg/l of octylphenol.

The daphnid lengths in the octylphenol mean measured concentrations of 0.062, 0.12 and 0.23 mg/l were significantly different from the pooled controls. Length measurements could not be made on the 0.51 mg/l test levels, since all daphnids had died by day 9.

Mean measured	Mean % Survival	Mean adult Mean you	Mean young/adult reproduction	
concentration (m	g/l)	length (mm)	day	
Control	98	3.80	5.0	
Solvent control	95	3.77	4.9	
Pooled control	96	3.79	5.0	
(Level 1) 0.037	100	3.74	4.9	
(Level 2) 0.062	95	3.69	4.6	
(Level 3) 0.12	100	3.64	3.4	
Level 4) 0.23	100	3.43	2.5	
(Level 5) 0.51	0*	*	*	

^{*} All daphnids had died by day 9 of the study.

CONCLUSIONS

A 21 day EC50 of 0.34 mg/l, based on growth and survival was determined.

The study accurately estimates the toxicity of octylphenol to *Daphnia magna*. Statistically the data identifies the MATC limits for the test substance as >0.037 and <0.062 mg/l. These limits represent the worst case estimate. Since the mean adult length of the 0.062 mg/l test concentrations is only 2.6% less than the pooled control, the statistical difference indicated may not be biologically significant. This is supported by the fact that the reproduction data did not identify an effect at this level. Therefore, a case can be made that based on biological effects, a more conservative MATC of >0.062 and <0.12 mg/l may be more appropriate.

DATA QUALITY

IUCLID 2000

GLP study

Method follows EPA/TSCA guideline.

Information taken from the test report.

REFERENCES

IUCLID 2000, SIDS 2000 and Waern 2000 as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

Also cited in the Targeted Environment Risk Assessment Report for 4-tert-octylphenol, Cas No. 140-66-9; Draft Environment Report, June 2001, UK Environment Agency.

Chronic Toxicity of Octylphenol to *Daphnia magna* Under Flow-Trough Test Conditions

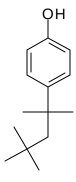
Study Number 36195

12.4 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- tert- Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.51 mg/l

ChV (21day) 0.032 mg/l Remark: log Kow used 5.28 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.51 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

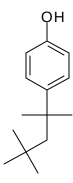
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

OECD Guideline No. 401

GLP: (Y)

Year study performed: 1991

Species: Sprague-Dawley strain rat.

RESULTS

LD50: >2000 mg/kg

CONCLUSIONS

The test substance has not been classified in the Acute Oral Toxicity Test.

LD50: >2000 mg/kg.

DATA QUALITY

Performed according to OECD guidelines and GLP.

Information taken from IUCLID database.

Purity of the test substance / decomposition: Information not available in database.

REFERENCES

IUCLID database

SafePharm Laboratories Ltd., P.O Box No. 45, Derby, DE1 2BT, U.K. phenol, (4-(1,1,3,3-tetramethylbutyl): range-finding acute oral toxicity test in the rat. Unpublished report project No. 47/1578, March 1991.

OTHER

Sandoz Chemikalien AG, Muttenz (CH).

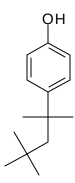
Huels AG Marl.

13.2 ACUTE TOXICITY - ADDITION ORAL

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

OECD Guideline No. 401

GLP: (Y)

Year study performed: 2000

Species: Rat.

RESULTS

LD50: 2200 mg/kg bw

CONCLUSIONS

LD50: 2200 mg/kg bw.

DATA QUALITY

Carried out to standard guidelines according to OECD.

Purity of the test substance / decomposition: Information not available in database.

REFERENCES

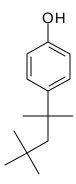
Waern (2000) as referenced in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced under contract for the UK Environment Agency by WRc-NSF, UK.

13.3 ACUTE TOXICITY INHALATION

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

Method/guideline followed: Information not available in database

GLP: (N)

Year study performed: 1973

Species: albino rat.

No of animals per dose: 6 adult rats.

Route of administration: Inhalation exposure.

Remarks: Test substance:

p-tert-octylphenol - 89%

o-tert-octylphenol - 2%

dioctylphenol - 5%

other isomers - 3%

phenol - 1%

RESULTS

Inhalation exposure of 6 adult albino rats at a concentration of 116 mg/l for one hour caused death of all animals within 24 hours:

Observations: Respiratory distress.

Gross autopsy: Pulmonary hemorrhage.

CONCLUSIONS

Inhalation exposure of 6 adult albino rats at a concentration of 116 mg/l for one hour caused death of all animals within 24 hours:

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

REFERENCES

IUCLID database

Rohm & Haas Co., 1973.

EPA/OPTS Public File 878213508, microfiche no.: 0205842.

OTHER

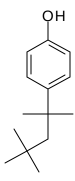
Source: Huels AG Marl.

13.4 ACUTE TOXICITY DERMAL

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

Method/guideline followed: Information not available in database

GLP: (N)

Year study performed: 1981

Species: Rabbit.

Remarks: Test substance: purity ca 95%.

RESULTS

LD50: >2000 mg/kg.

CONCLUSIONS

The test substance has not been classified in the Acute Dermal Toxicity Test.

LD50: >2000 mg/kg.

DATA QUALITY

Not a GLP study. Information taken from IUCLID database

REFERENCES

IUCLID database. BASF AG, dept of toxicology, unpublished data, (79/298), 01-12-81.

OTHER

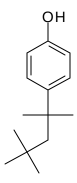
Source: BASF AG Ludwigshafen.

13.5 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

OECD Guidelines No. 404

GLP: (Y)

Year study performed: 1991

Species: New Zealand White rabbits.

Sex: 2 males, 1 female.

RESULTS

Erythema/Eschar formation: total scores after:

24 h = 372 h = 2

Oedema formation: total scores after:

24 h = 2

72 h = 0

The test material produced a primary irritation index of 1.2 and was classified as a mild irritant to rabbit skin according to the Draize classification scheme. No corrosive effects were noted.

CONCLUSIONS

Classified as a mild irritant to rabbit skin according to the Draize classification scheme and a non-irritant according to EEC labelling regulations.

DATA QUALITY

Performed according to OECD guidelines and GLP.

Information taken from IUCLID database and test report.

Purity of the test substance/decomposition: information not available in database.

REFERENCES

SafePharm Laboratories., P.O Box No. 45, Derby, DE1 2BT, U.K. acute dermal irritation test in the rabbit. Unpublished report, project No. 47/1579, Feb 1991 as cited in the IUCLID database.

OTHER

SIDS Dossier on the HPV P3 Chemical 140-66-9 Phenol, 4-(1,1,3,3-tetramethylbutyl)-, June 1993, received from Sandoz Chemikalien AG, Muttenz (CH).

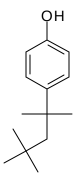
Huels AG Marl.

13.6 ACUTE TOXICITY EYE IRRITATION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

OECD Guidelines No. 405	
GLP: (Y)	
Year study performed: 1991	

Species: New Zealand White Rabbits.

RESULTS

A single application of the test material to the non-irrigated eye of one rabbit produced areas of translucent corneal opacity, iridial inflammation and severe conjunctival irritation. Other adverse ocular reactions noted were white appearance of the nictitating membrane, sloughing of the conjunctival surface, hemorrhage of the conjunctival membrane and blood stained discharge.

CONCLUSIONS

The test material was considered to be at least a severe irritant (Class 6 on a 1 to 8 scale) to the rabbit eye according to a modified Kay and Calandra classification system (based on one rabbit only). The test material was also considered to be an irritant according to EEC labelling regulations.

DATA QUALITY

Performed according to OECD guidelines and GLP.

Information taken from IUCLID database.

Purity of the test substance/decomposition: information not available in database.

REFERENCES

SafePharm Laboratories Ltd., P.O Box No 45, Derby, DE1 2BT, U.K. Phenol, 4-(1,1,3,3-tetramethylbutyl): acute eye irritation test in the rabbit. Unpublished report, project No. 47/1580, Feb 1991 as cited in the IUCLID database.

OTHER

SIDS Dossier on the HPV P3 Chemical 140-66-9 Phenol, 4-(1,1,3,3-tetramethylbutyl)-, June 1993, received from Sandoz Chemikalien AG, Muttenz (CH).

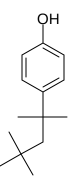
Huels AG Marl.

13.7 ACUTE TOXICITY SKIN SENSITISATION

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

OECD Guidelines No. 406

GLP: (N)

Year study performed: 1988

Species: Guinea-pig.

Remarks: Test substance = p-substituted octylphenol.

Challenge concentration: 20% in corn oil.

RESULTS

0/20 animals showed sensitisation 24 or 48 hours after the patch test.

CONCLUSIONS

The test material is not classified as a sensitizer.

DATA QUALITY

Performed according to OECD guidelines. Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance/decomposition: Information not available in database.

REFERENCES

IUCLID database

Huels report No. 1197, 1988 (unpublished).

OTHER

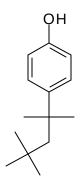
Source: Huels AG Marl.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATION)

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



METHOD

Method: Directive 84/449/EEC B.14 "Other effects – Mutagenicity(*Salmonella typhimurium* – reverse mutation assay)

Type: Reverse mutation assay

System of testing: Bacterial Plate incorporation

GLP: (Y)

Year study performed: 1984

Species/strain: Salmonella typhimurium TA 98, TA 100, TA 1537, TA 1538.

Metabolic activation: S9 mix (with and without).

Concentrations tested: $0 - 5000 \mu g/plate$.

RESULTS

The test substance did not induce any significant mutations in the test system.

CONCLUSIONS

The test substance is considered to be non mutagenic.

DATA QUALITY

Performed according to EU guidelines. GLP study.

Information taken from IUCLID database.

Purity of the test substance/decomposition: Information not available in database.

REFERENCES

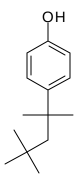
IUCLID database. Huels Report No. AM-91/11, 1991 (unpublished).

16.1 REPEATED DOSE TOXICITY 3 MONTHS DIETARY

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

Method/guideline followed: Information not available in database, similar to EPA Health Effects Testing Guidelines

GLP: (N)

Year study performed: 1982

Species/strain: Rat BOR/WISW (SPF Cpb).

No. of rats per sex per dose: 10 rats/sex/dose.

Route of administration: Oral feed.

Sex: Males and females.

Exposure Period: 3 months.

Dose/Concentration: 0, 30, 300 and 3000 ppm.

Frequency of treatment: Daily.

Control group and treatment: Yes, concurrent no treatment.

Post exposure observation period: None.

Test Conditions: Twenty rats (10 females and 10 males) were used for the control and each of the three treatment groups. The animals were exposed to the test substance for 90 days. Treatment effects were determined by statistical comparison of mortality, body weight changes, food and water consumption, organ weight changes, clinical chemistry, hematology and histological evaluation of tissues from sacrificed animals. Although the study was conducted prior to the issuance of EPA's Health Effects Testing Guidelines, it closely follows the guidelines. The test procedures, including animal and dose

selection, exposure conditions, observations, test substance administration, hematology, clinical chemistry, urinalysis and gross necropsy were in accordance with the testing guidelines.

RESULTS

NOAEL = 30 ppm.

LOAEL = 300 ppm.

Histologic examinations were conducted on only five animals per sex from the high dosage group as opposed to the ten animals per sex suggested in the EPA guidelines. However, the lack of a treatment-related effect in the high exposure group strongly suggests that treatment-related effects would not have been observed if all of the animals had been evaluated.

Throughout the study, no clinical signs of toxicity were observed. Furthermore, no significant reduction in food consumption was observed in either sex at any dosage level. A 28% increase in water consumption was noted in female rats exposed to the high dosage level. However, this effect was not observed in the male rats at any dosage level.

In both sexes, mean body weight gain was significantly reduced in the high dose animals. In females exposed at the high dosage, the heart weight was decreased and the brain weight was increased. The kidney, testes and brain weight were increased in males exposed at the high dosage level. In addition, males exposed to 300 ppm of the test substance experienced an increase in brain weight. Relative brain weight was the only dose –related organ weight change that was statistically significant.

Hematologic parameters in all treated male rats were unaffected by exposure to the test substance. A decrease in hemoglobin and hematocrit was observed among female rats in the high dosage group. Although thyroxin concentration in female rats at the high dose level was increased after one month, thyroxin concentration was not significantly elevated following three months of exposure. The increase at one month was attributed to elevated values in two female rats. Since no increase in thyroxin values was noted at three months and no histopathologic findings were observed in the thyroid gland, the increased thyroxin values were not considered to be toxicologically significant. All other findings were either not significantly different from controls, not dosage-related, or were within the normal range for animals in this age group.

At the gross necropsy no treatment-related abnormalities were observed. In addition, no treatment-related histopathologic observations were noted.

CONCLUSIONS

NOAEL = 30 ppm.

The results from this study demonstrate that high level exposure to the test substance produces low level toxic effects.

DATA QUALITY

Not a GLP study.

Information taken from the IUCLID database and report.

Test substance p-substituted octyl (purity 93.1%).

REFERENCES

IUCLID

Bayer report No. 10733, 1982 EPA/OPTS Public file 878213507, microfiche no.: 0205841.

OTHER

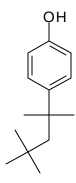
Source: Huels AG Marl.

16.2 REPEATED DOSE TOXICITY 3 MONTHS DIETARY

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

Method/guideline followed: Information not available in database

GLP: (N)

Year study performed: 1961

Species/strain: Rat Albino.

No. of rats per sex per dose: 15/sex/dose.

Route of administration: Oral feed.

Sex: Males and females.

Exposure Period: 3 months.

Dose/Concentration: 5% in diet.

Frequency of treatment: Daily.

Control group: Yes.

Post exposure observation period: None.

Test Conditions: Fifteen young male and young female albino rats were placed on the following dietary concentrations of the test substance: 0 and 5%. Finely ground Purina Dog Chow Kibbled Meal served as the basic diet and into this was incorporated a 35% aqueous solution of the test substance in amount calculated to achieve a 5% concentration of the active ingredient. An equivalent amount of water was added to the diet of the control rats. The animals were individually caged and were weighed once a week. Food consumption data were collected over a 3-day period during the thirteenth week. Urine collected during the last week from 5 rats of each sex at each dietary level was tested semi-quantitatively for glucose (Morris Anthrone method) and protein (Pro-Teen, sulfosalicyclic acid and

Shevky & Stafford methods). Hematologic determinations on 5 rats of each sex at each dietary level were made at the end of the test period.

Organ to body weight ratios for liver, kidney, spleen, heart and testes were determined at sacrifice of the 3-month survivors. Tissues taken for histopathologic study were: heart, lung, liver, kidney, spleen, gastroenteric, bladder, bone marrow, muscle, skin, brain, thyroid, adrenal and pancreas.

RESULTS

No effects of treatment were apparent and there were no deviations in food consumption. Urinary concentrations of glucose and protein were comparable in treated and control animals. There were no hematologic related effects to treatment. There were no statistically related differences for organ to body weight between control and treated animals. There were no lesions apparent.

CONCLUSIONS

Feeding of 5% of the test substance in the diet of rats for a period of 3 months had no effect on growth, survival, food consumption, urinary excretion of glucose and protein, hematologic values, or organ to body weight ratios, and produced no pathologic lesions.

DATA QUALITY

Not a GLP study.

Information taken from the IUCLID database and report.

REFERENCES

IUCLID database.

Rohm and Haas, 1961 EPA/OPTS Public file 40-6162030, microfiche no.: 0527757.

OTHER

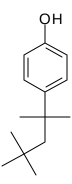
Source: Huels AG Marl.

17.1 TOXICITY TO REPRODUCTION

TEST SUBSTANCE

p-tert-Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

Guideline followed: OPPTS Draft Testing Guidelines (870.3800; 1996 with additions)

Type: Two generation Reproduction study (one litter per generation)

GLP: unconfirmed

Year study performed: unknown, but after 1996

Species: Rat/CD® (Sprague-Dawley)

Route of administration: Oral diet

Doses levels: (dose range selected to evaluate potential toxicity at conventional high dosage, and also to address the possibility of low-dose estrogenic effects)

0, 0.2, 20, 200, and 2000 ppm = 0, 0.01, 1, 10 and 100 mg/kg bodyweight/day- male adults

0, 0.2, 20, 200, and 2000 ppm = 0, 0.015, 1.5, 15 and 150 mg/kg bodyweight/day- female adults

0, 0.2, 20, 200, and 2000 ppm = 0, 0.08, 8, 80 and 800 mg/kg bodyweight/day- weanlings*

Male adult bodyweight 600g; daily average food intake 30 g

Female adult bodyweight 400g; daily average food intake 30 g

Weanlings average bodyweight 50g; daily average food intake 20 g

Number of animals and sex: 30 male and 30 female per group.

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Age of animals before initiation of dosing: not known

Frequency of treatment: Continuous from study day 1 until necropsy/termination

Pre-mating exposure for males and females: 10 weeks

Study design:

F0 and F1 Animals were randomly mated within treatment groups for a two-week mating period to produce the next generation. Males were necropsied after the delivery period. Litters were culled to 10 pups on postnatal day (pnd) 4 and weaned on pnd 21.

At weaning, up to 3 weanlings/sex/litter were necropsied, and 30/sex/dose were selected as parents of the next generation.

For F1 weanlings, acquisition of vaginal patency in females and preputial separation in males was assessed, and vaginal cytology for estrous cyclicity was evaluated for three weeks prior to mating.

On the day of birth, anogenital distance was measured for F2 offspring. F2 weanling females were retained through acquisition of vaginal patency, at which time they were terminated without further evaluation. F2 males were retained through acquisition of preputial separation, until age 111±5 days, when they were subjected to necropsy.

Parental females were necropsied after weaning of their litters, with organ weights and histopathology including enumeration of ovarian follicles.

Parental males were necropsied after the littering period, with reproductive organ weights, epididymal sperm count, sperm motility and morphology, testicular homogenization-resistant spermatid count, and histopathology. In addition, dorsal prostate weight and epididymal sperm and testicular spermatid measurements were recorded for the F2 males.

Statistical methods:

Not known

RESULTS

F0 results

Adult systemic toxicity was evident for F0 parental animals at 2000 ppm, expressed as consistent and persistent reductions in body weights and weight gains during the pre-mating period.

Feed consumption in g/day and g/kg/day and food efficiency was variable.

There were no treatment or dose-related clinical signs of toxicity in either sex.

Body weights during gestation were unaffected and were reduced during lactation in females at 2000 ppm.

There were no effects of treatment in F0 females on mating, fertility, pregnancy, or gestational indices.

There were also no effects of treatment in males on mating or fertility indices.

Estrous cycle length and stage of estrous at necropsy were equivalent across all F0 females.

Paired ovarian follicle counts were equivalent between high dose and control groups for F0 females.

Gestational length in days was equivalent across all groups for females.

Necropsy: In parental male rats absolute organ weights were almost uniformly unaffected for the liver, kidneys, adrenal glands, spleen and brain. Relative organ weights exhibited only occasional increases, almost exclusively at 2000 ppm, most likely due to the reduced body weights at this dietary concentration. There were no treatment-related gross or microscopic findings on these organs.

Absolute and relative uterine weights were significantly reduced at 2000 ppm; ovarian weights were unaffected.

There were no effects of treatment on any adult male reproductive organs, including no effects on absolute or relative weights of testes, epididymides, prostate, dorsal prostate, seminal vesicles with coagulating glands, and no gross or microscopic effects of treatment on these organs. There were also no effects of treatment on epididymal sperm concentration, percent motile or progressively motile sperm, testicular homogenisation-resistant spermatid head counts, daily sperm production, efficiency of daily sperm production, or percent abnormal sperm for males.

F1 results

Adult systemic toxicity was evident for F1 parental animals at 2000 ppm, expressed as consistent and persistent reductions in body weights and weight gains during the prebreed.

Feed consumption in g/day and g/kg/day and food efficiency was variable.

There were no treatment or dose-related clinical signs of toxicity in either sex.

Body weights during gestation were unaffected and were reduced during lactation in females at 2000 ppm.

There were no effects of treatment in F1 females on mating, fertility, pregnancy, or gestational indices.

There were also no effects of treatment in males on mating or fertility indices.

Estrous cycle length and stage of estrous at necropsy were equivalent across all F1 females.

Paired ovarian follicle counts were equivalent between high dose and control groups for F1 females.

Gestational length in days was slightly but statistically significantly prolonged at 0.2 ppm (by 0.3 days). Gestation length was unaffected at 20, 200 or 2000 ppm for F1 females.

For F1 offspring, there were no effects of treatment on stillbirth or live birth indices, for number of live pups per litter on pnd 0, sex ratio (% males) throughout lactation, lactational survival index (pnd 4-21), or 4-, 7-, 14- or 21-day survival indices.

Pup body weights per litter were significantly reduced at 2000 ppm on PND 14 and 21.

Reduced pup weights were observed only during the latter portion of the lactational period when the pups generally began to self-feed and therefore were likely to be directly exposed to the test chemical in the diet (CD® rat pups begin to self-feed late in the 2^{nd} week of life).

Acquisition of vaginal patency and preputial separation in offspring was significantly delayed (by less than two days) at 2000 ppm, most likely related to the lower body weights of these pups at this dose.

These minimal effects on reproductive development in the offspring required (by guideline) measurement of anogenital distance in F2 offspring at birth (pnd 0)

Necropsy:

In parental male rats absolute organ weights were almost uniformly unaffected for the liver, kidneys, adrenal glands, spleen and brain. Relative organ weights exhibited only occasional increases, almost exclusively at 2000 ppm, most likely due to the reduced body weights at this dietary concentration. There were no treatment-related gross or microscopic findings on these organs. Absolute and relative weights of F1 female reproductive organs (ovaries and uterus) were unaffected by treatment.

There were no effects of treatment on any adult male reproductive organs, including no effects on absolute or relative weights of testes, epididymides, prostate, dorsal prostate, seminal vesicles with coagulating glands, and no gross or microscopic effects of treatment on these organs. There were also no effects of treatment on epididymal sperm concentration, percent motile or progressively motile sperm, testicular homogenisation-resistant spermatid head counts, daily sperm production, efficiency of daily sperm production, or percent abnormal sperm for males.

F2 results

Adult systemic toxicity was evident for F2 retained animals at 2000 ppm, expressed as consistent and persistent reductions in body weights and weight gains during the postwean exposure period.

Feed consumption in g/day and g/kg/day and food efficiency was variable.

There were no treatment or dose-related clinical signs of toxicity in either sex.

For F2 offspring, there were no effects of treatment on stillbirth or live birth indices, for number of live pups per litter on pnd 0, sex ratio (% males) throughout lactation, lactational survival index (pnd 4-21), or 4-, 7-, 14- or 21-day survival indices.

Pup body weights per litter were significantly reduced at 2000 ppm on PND 14 and 21.

Reduced pup weights were observed only during the latter portion of the lactational period when the pups generally began to self-feed and therefore were likely to be directly exposed to the test chemical in the diet.

Acquisition of vaginal patency and preputial separation in offspring was significantly delayed (by less than two days) at 2000 ppm, most likely related to the lower body weights of these pups at this dose.

F2 male pups exhibited no effect of treatment on anogenital distance. F2 female pups exhibited statistically significantly longer mean anogenital distances at 0.2 ppm (0.79 mm), 20 ppm (0.81 mm), 200 ppm (0.85 mm), and 2000 ppm (0.79 mm) relative to the control group value (0.76 mm); these changes were not considered biologically significant.

Necropsy:

In retained F2 male rats absolute organ weights were almost uniformly unaffected for the liver, kidneys, adrenal glands, spleen and brain. Relative organ weights exhibited only occasional increases, almost exclusively at 2000 ppm, most likely due to the reduced body weights at this dietary concentration. There were no treatment-related gross or microscopic findings on these organs.

There were no effects of treatment on any adult male reproductive organs, including no effects on absolute or relative weights of testes, epididymides, prostate, dorsal prostate, seminal vesicles with coagulating glands, and no gross or microscopic effects of treatment on these organs. There were also no effects of treatment on epididymal sperm concentration, percent motile or progressively motile sperm, testicular homogenisation-resistant spermatid head counts, daily sperm production, efficiency of daily sperm production, or percent abnormal sperm for F2 retained adult males.

CONCLUSION

NOAELs for systemic and postnatal toxicity were 200 ppm and at or above 2000 ppm for reproductive toxicity.

Dietary exposure to octylphenol for two generations, one litter per generation, at 0, 0.2, 20, 200 and 2000 ppm, resulted in:

Decreased body weights and weight gains at 2000 ppm

Offspring toxicity (reduced body weight during lactation) at 2000 ppm

Delayed vaginal opening and preputial separation at 2000 ppm considered related to body weight decreases

No effects on reproductive parameters

No effects on testes weights or morphology

No effects on epididymal sperm counts, motility or morphology

No effects on daily sperm production or efficiency of daily sperm production

No estrogen-like effects on males or females

No effects on prostate or dorsal prostate weights or histopathology.

DATA QUALITY

Study performed using OPPTS Draft Testing Guidelines (870.3800;1996 with additions)

REFERENCES

Two-Generation Reproductive Toxicity Evaluation of Para-tert Octylphenol Administered in the Feed to CD® (Sprague-Dawley) Rats (Abstract)

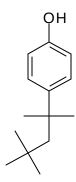
Sponsor: Union Carbide Corporation, Health, Safety & Environmental, 39 Old Ridgebury Road, Danbury, CT 06817-0001, USA.

17.2 TOXICITY TO REPRODUCTION (ESTROGENIC ACTIVITY) - ADDITION

TEST SUBSTANCE

*p-tert-*Octylphenol

CAS No. 140-66-9



IUCLID Identification Number: 140-66-9

METHOD

Guideline followed: E Screen/ MCF7/ Yeast Screen

GLP: No

Year study performed: Around 2000

RESULTS

E Screen Relative Potency = 0.03

MCF7 Relative Potency = 0.2

Yeast Screen Relative Potency = 0.07

Potency Relative to estradiol = 100

CONCLUSION

E Screen Relative Potency = 0.03

MCF7 Relative Potency = 0.2

Yeast Screen Relative Potency = 0.07

Potency Relative to estradiol = 100

DATA QUALITY

Included in UK Initial Assessment.

REFERENCES

Waern (2000) as summarised in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

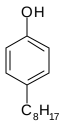
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Melting point: 82.77℃ (weighted value).

CONCLUSIONS

The test substance has a calculated melting point of 82.77°C.

DATA QUALITY

Calculation method used.

REFERENCES

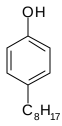
SYRACUSE Chemical Properties Prediction Program. MPBPWIN v1.30.

2.1 BOILING POINT

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): No

Year study performed: 1992.

RESULTS

Boiling point: 296°C at 760 mm Hg without decomposition.

CONCLUSIONS

The test substance has a boiling point of 296°C at 760 mm Hg.

DATA QUALITY

Not a GLP study.

Information taken from a technical datasheet.

Purity of the test substance: 85.0% octylphenol min. and 15.0% nonylphenol max.

REFERENCES

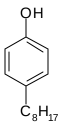
Schenectady International Inc. Technical Datasheet (7/92).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 0.000098 mm Hg @ 25°C (0.013 Pa).

CONCLUSIONS

The test substance has a vapor pressure of 0.000098 mm Hg @ 25°C.

DATA QUALITY

Calculation method.

REFERENCES

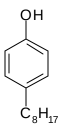
SYRACUSE Chemical Properties Prediction Program. MPBPWIN v1.30.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 5.50

CONCLUSIONS

The test substance has a log Kow of 5.50.

DAT A QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v1.63.

OTHER

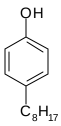
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 3.114 mg/l at 25℃.

CONCLUSIONS

The solubility of the test substance in water is 3.114 mg/l at 25°C.

DATA QUALITY

Calculation method used.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v1.33.

OTHER

Log Kow value used: 5.50.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

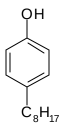
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $50.3 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.55 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.55 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

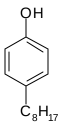
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 206
Data temperature (°C): 25
Water solubility (mg/l): 3.114
Vapor pressure (Pa): 0.013
Log Kow: 5.50
Melting point (°C): 83

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	0.060
Soil	97.4
Water	0.35
suspended sediment	0.068
fish	0.0055
Sediment	2.16

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

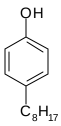
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Primary biodegradation: in days.

Ultimate biodegradation: in weeks.

CONCLUSIONS

The program predicts that under the stated conditions the test substance will biodegrade rapidly. This material is considered to be inherently biodegradable, and would aerobically biodegrade.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

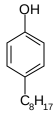
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.21 mg/l

ChV(30 day) 0.030 mg/l

ChV (90 day) 0.007 mg/l

Remark: log Kow used 5.50 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.21 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

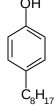
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.082 mg/l

ChV (96hr) 0.062 mg/l

Remark: log Kow used 5.50 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.082 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

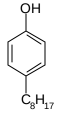
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.41 mg/l

ChV (21day) 0.023 mg/l

Remark: log Kow used 5.50 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.41 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

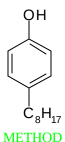
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

p-Octylphenol

CAS No. 1806-26-4



EPA 40 CFR 163.81-1 Defined Oral LD₅₀

GLP: (N)

Year study performed: 1985

Species: Albino rats

Sex: Males and Females

No of animals per sex per dose: 5 males and 5 females

Vehicle: Corn oil

Route of administration: Oral, gavage

Remarks: Following a range finding study, 5 groups of ten fasted animals were given a single oral dose of the test material. Dose range used was 0.25-2.0 g/kg. After dosing the animals were observed for 14 days.

RESULTS

LD50 Total 1.2 g/kg, Male 1.4 g/kg, Female 1.1 g/kg

95% confidence limits: Upper 1.78 g/kg, Lower 0.9 g/kg

Dosage 0.25 g/kg and 0.5 g/kg: no signs of toxicity.

Dosage 1.0, 1.5 and 2.0 g/kg: increasing signs of toxicity with dosage with death of all animals at 2.0 g/kg.

Clinical signs included diarrhoea, lethargy, inability to move, bloody nasal discharge chromodacryorrhoea and loss of appetite.

CONCLUSIONS

LD50: 1.2 g/kg

EPA Category III

DATA QUALITY

No evidence of GLP. Conducted to EPA guidelines.

REFERENCES

Product Safety Labs, 725 Cranbury Road, East Brunswick, New Jersey 08816, USA study report no. T-5049

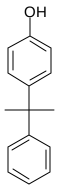
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1992.

RESULTS

Freeze point: 72°C.

CONCLUSIONS

The freezing point of the test substance is 72%.

DATA QUALITY

Not a GLP study.

Information taken from technical datasheet.

Purity given as 99% minimum.

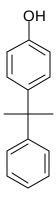
REFERENCES

Schenectady International Inc. Technical datasheet (6/92).

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1992.

RESULTS

Boiling point: 335°C at 760 mm Hg.

CONCLUSIONS

The boiling point of the test substance is 335°C at 760 mm Hg.

DATA QUALITY

Not a GLP study.

Information taken from technical datasheet.

Decomposition: information not available.

Purity given as 99% minimum.

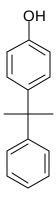
REFERENCES

Schenectady International Inc. Technical datasheet (6/92)

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 2.28×10^{-5} mm Hg at 25 °C (0.0030 Pa).

CONCLUSIONS

The calculated vapor pressure is 2.28×10^{-5} mm Hg at 25° C.

DATA QUALITY

Calculation.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBWIN v 1.30.

OTHER

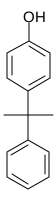
Modified Grain Method used.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 4.12.

CONCLUSIONS

The log Kow of the test substance is 4.12.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63.

OTHER

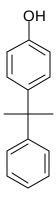
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: calculation method using a log Kow of 4.12.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 43.27 mg/l at $25 ^{\circ}\text{C}$.

CONCLUSIONS

The calculated water solubility is 43.27 mg/l at 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

The log Kow value obtained using the programme KOWWIN was used in the calculation of water solubility (see 4.1).

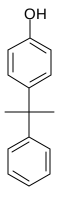
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $44.8 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.87 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.87 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

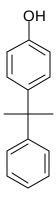
No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

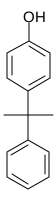
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:212Data temperature (°C):25Water solubility (mg/l):43.27Vapor pressure (Pa):0.0030Log Kow:4.12Melting point (°C):72

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	0.023
Soil	90.2
Water	7.72
suspended sediment	0.063
fish	0.0051
Sediment	2.00

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

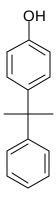
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days/weeks

Ultimate biodegradation in weeks/months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

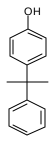
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p- (alpha, alpha-Dimethylbenzyl) phenol

CAS No. 599-64-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 1.54 mg/l

ChV(30 day) 0.22mg/l

ChV (90 day) 0.029 mg/l

Remark: log Kow used 4.12 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 1.54 mg/l.

DATA QUALITY

Estimated

REFERENCES

ECOSAR v.0.99e

OTHER

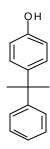
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p- (alpha, alpha- Dimethylbenzyl) phenol

CAS No. 599-64-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 1.40 mg/l

ChV (96hr) 0.48 mg/l

Remark: log Kow used 4.12 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 1.40 mg/l.

DATA QUALITY

Estimate

REFERENCES

ECOSAR v.0.99e

OTHER

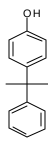
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p- (alpha, alpha- Dimethylbenzyl) phenol

CAS No. 599-64-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 1.6 mg/l

ChV (21day) 0.17 mg/l

Remarks: log Kow used 4.12 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 1.6 mg/l.

DATA QUALITY

Estimated

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

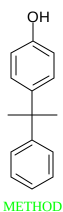
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



EPA TSCA 40 CFR 798.1175

GLP: (Y)

Year study performed: 1990

Species: Sprague-Dawley rats, young adult

Sex: Males and females

No of animals per sex per dose: Five groups of male and female rats

Vehicle: The test material was prepared as a 25% w/v formulation in corn oil

Route of administration: Oral

Remarks: Four dose levels were evaluated: 5.0, 2.5, 1.25 and 0.625 g/kg. Animals were observed for 14 days.

RESULTS

Clinical signs:

Brownish red and saliva like stains on muzzle, faecal stains, piloerection, unkempt fur, moderate and/or slight depression, extensive yellow brown stains on anal and stomach area, red stains around eyes and hunched posture were noted in the two highest dose levels.

Red stains on muzzle, piloerection, slight depression, faecal and urine stains, brown yellow stains on urogenital area, rapid breathing and unkempt fur were seen in animals dosed at 1.25 or 0.625 g/kg

Number of deaths at each dose level:

At the dose level of 5.0 g/kg four deaths (100%), were noted between days one and two of the observation period.

At the dose level of 2.5 g/kg four deaths (100%), were noted between days one and five of the observation period.

At the dose level of 1.25 g/kg no deaths (0%), were noted during the observation period.

At the dose level of 0.625 g/kg no deaths (0%), were noted during the observation period.

CONCLUSIONS

The test material is classified in Toxicity Category III (greater than 500 mg/kg - 5000 mg/kg) by oral administration.

DATA QUALITY

Study conducted to EPA guidelines and GLP.

REFERENCES

Study conducted by Hill Top Biolabs Inc.

OTHER

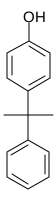
Unpublished study report for Schenectady International, Inc. Schenectady, NY "Acute Oral Toxicity in Rats-p-cumylphenol- Median Lethal Dosage Determination. August 23, 1990".

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

p-(alpha, alpha-Dimethylbenzyl)phenol

CAS No. 599-64-4



METHOD

Type: The Salmonella Reverse Mutation Assay

GLP: Not known

Year study performed: 1984

Species/strain: Salmonella typhimurium TA 98, TA 100, TA 1535 and TA 1537

Metabolic activation: S9 fraction from male SD strain rats

Concentrations tested: 5, 10, 50, 100 and 500 μ g/plate

RESULTS

>100 µg/plate demonstrated cytotoxicity in strains with metabolic activation.

>50 µg/plate demonstrated cytotoxicity in strains without metabolic activation.

The results of the test were negative for mutagenic effects in the presence or absence of metabolic activation.

CONCLUSIONS

The test substance is not genetically active in The Salmonella Reverse Mutation Assay.

DATA QUALITY

Not to GLP

REFERENCES

Report on Mutagenicity test using microorganisms

OTHER

Unpublished study report performed by Mitsui Petrochemical Industries for General Electric Company; December 12, 1984. No further information available.

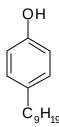
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: based on USEPA TSCA Environmental Fate Test Guidelines (USEPA 1985). Measurement of Crystallisation Point.

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Crystallisation point: 24.5°C

CONCLUSIONS

The test substance has a crystallisation point of 24.5°C.

DATA QUALITY

GLP study.

Information taken from IUCLID database.

Purity of the test substance given as > 95% p-nonylphenol, confirmed by subsequent gas chromatography analysis.

REFERENCES

Alkylphenol and Ethoxylates Panel (1990). Five physical/chemical 4-nonylphenol final reports. 1. boiling point. 2. crystallisation point (instead of melting point. 3. dissociation constant. 4. water solubility. 5. vapor pressure. Chemical Manufacturers Association, Washington DC, 20037, August 21, 1990 as cited in IUCLID database.

2.1 BOILING POINT

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3

METHOD

Method/guideline followed: US EPA (1989, 40 CFR 796.1220) with modifications for the high temperatures required.

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Three initial trials were run. In all cases the test substance decomposed before the boiling point was reached at temperatures ranging from 215 to 233 (mean 230) °C. In a subsequent trial, decomposition of the test material was observed at temperatures ranging from 283 to 295°C. The boiling point was reported as being greater than 300 °C. However, data from this study indicate that the test substance will thermally decompose before boiling.

CONCLUSIONS

The test substance decomposes before the boiling is reached.

DATA QUALITY

GLP study.

Information taken from IUCLID database.

Purity of the test substance given as > 95% p-nonylphenol, confirmed by subsequent gas chromatography analysis.

REFERENCES

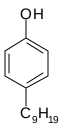
Alkylphenol and Ethoxylates Panel (1990). Five physical/chemical 4-nonylphenol final reports. 1. boiling point. 2. crystallisation point (instead of melting point. 3. dissociation constant. 4. water solubility. 5. vapor pressure. Chemical Manufacturers Association, Washington DC, 20037, August 21, 1990 as cited in IUCLID database.

2.2 BOILING POINT

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1992.

RESULTS

Boiling point: 310°C @ 760 mm Hg.

CONCLUSIONS

The test substance has a boiling point of 310°C @ 760 mm Hg.

DATA QUALITY

Not a GLP study.

Information taken from technical datasheet.

Decomposition: information not available.

Purity of the test substance given as typically 95.0% minimum.

REFERENCES

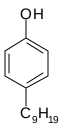
Schenectady International Inc. Technical Datasheet (7/92).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: US EPA 40 CFR 795.1950 – Gas saturation apparatus method

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Vapor pressure: ca. 0.00455 Pa @ 25°C.

CONCLUSIONS

The test substance has a vapor pressure of ca. 0.00455 Pa @ 25° C.

DATA QUALITY

GLP study.

Information taken from IUCLID database.

Decomposition: information not available.

Purity of the test substance given as > 95% *p*-nonylphenol, confirmed by subsequent gas chromatography analysis.

REFERENCES

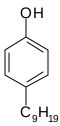
Alkylphenol and Ethoxylates Panel (1990). Five physical/chemical 4-nonylphenol final reports. 1. boiling point. 2. crystallisation point (instead of melting point. 3. dissociation constant. 4. water solubility. 5. vapor pressure. Chemical Manufacturers Association, Washington DC, 20037, August 21, 1990 as cited in IUCLID database.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: the octanol-water partition coefficient (Kow) of nonylphenol was determined at two concentrations in accordance with USEPA guidelines. test vessels (25 ml Teflon centrifuge tubes) contained 18 ml of pH buffer, 1.9 ml of n-octanol and 100 μ l of a stock solution of 4-nonylphenol in n-octanol. Test vessels were agitated for one hour at 25°C and centrifuged at 10000 g for 30 minutes. The test substance was quantified in samples of octanol and water from each vessel by HPLC. The test substance was below the detection limit (32.5 μ g/l) in all water samples. Therefore, Kow values were reported as "greater than" values.

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Log Kow 3.8 – 4.77 @ 25°C

The data showed concentration dependence because the test substance was non-detectable in all water samples and the value $< 32.5~\mu g/l$ was used to calculate Kow.

CONCLUSIONS

The test substance has a log Kow of $3.8 - 4.77 @ 25^{\circ}C$.

DATA QUALITY

GLP study.

Purity of the test substance given as > 95% *p*-nonylphenol, confirmed by subsequent gas chromatography analysis.

Method of analysis was insufficiently sensitive to detect analyte in the aqueous phase. The results therefore probably underestimate the value of log Kow.

REFERENCES

Chemical Manufacturers Association (1991). Determination of the Octanol/Water Partition Coefficient of 4-Nonylphenol. Testing laboratory: RF Waston Inc. Lionville, PA. Study no. 90-046. Test sponsor: Chemical Manufacturers Association, Washington DC. December 1991.

OTHER

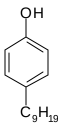
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

4.2 PARTITION COEFFICIENT

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: OECD Guide-line 107 "Partition Coefficient (n-octanol/water), Flask-shaking method.

GLP (Y/N): no.

Year study performed: 1989.

RESULTS

Log Kow: 3.28 @ 20°C.

CONCLUSIONS

The test substance has a log Kow 3.28 @ 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

The flask-shaking method is an acceptable technique for determining partition coefficients.

Purity of the test substance: information not available in database.

REFERENCES

Huels study 1989 (unpublished). Cited in IUCLID database.

OTHER

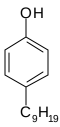
Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: USEPA TSCA Environmental Fate Test Guidelines

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Water solubility: 3.93 mg/l @ 25°C.

Remarks: The seawater solubility value was calculated as the mean dissolved 4-nonylphenol concentration in the three test samples following HPLC analysis of artificial seawater flowing over a column packed with nonylphenol. The solubility of 4-nonylphenol in the artificial seawater was determined to be 3.93 mg/l (standard deviation 0.38 mg/l, %RSD 10.5)

CONCLUSIONS

The solubility of the test substance in seawater is 3.93 mg/l @ 25°C.

DATA QUALITY

GPL study.

Information taken from IUCLID database.

Purity of the test substance given as > 95% p-nonylphenol, confirmed by subsequent gas chromatography analysis.

REFERENCES

ICI Chemicals & Polymers Limited Runcorn, Cheshire

HUELS AG MARL, 1990 as cited in IUCLID dataset.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

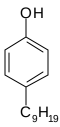
The solubility reported in this robust summary is for seawater therefore a calculated value has also been included (see 5.2).

5.2 WATER SOLUBILITY

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: calculation using a log Kow of 5.92.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 1.157 mg/l @ 25℃.

CONCLUSIONS

The test substance has a calculated water solubility of 1.157 mg/l @ 25°C.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Log Kow value of 5.92 used (as calculated by the programme KOWWIN).

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

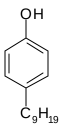
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Over all hydroxyl rate constant: $51.7 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.48 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.48 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

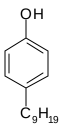
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 220
Data temperature (°C): 25
Water solubility (mg/l): 3.93
Vapor pressure (Pa): 0.0046
Log Kow: 3.28
Melting point (°C): 24.5

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	1.87
Soil	60.7
Water	36.0
suspended sediment	0.042
fish	0.0034
Sediment	1.35

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

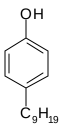
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: ISO Draft "BOD test for insoluble substances".

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1990.

Contact time: 28 days.

Inoculum: non-adapted, domestic, activated sludge.

Concentration: 34 mg/l.

RESULTS

Percentage degradation: 7% after 28 days.

CONCLUSIONS

p-Nonylphenol was not readily biodegradable under the test conditions.

DATA QUALITY

The reviewer for the IUCLID database considered this study to be valid without restriction and comparable to a guideline study.

REFERENCES

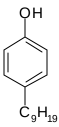
Huels AG: Report no. BO-90/3, 1990 (unpublished), as cited in IUCLID database.

9.2 BIODEGRADATION - ADDITION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



Commercial grade highly branched alkyl chain: 95% p-nonylphenol.

METHOD

Method/guideline followed: OECD 301 B, CO₂ Evolution, Modified Stürm Test

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1996.

Contact time: 28 days.

Inoculum: non-adapted, domestic, activated sludge.

Concentration: 12.2 mg/l (10 mg Carbon/l).

RESULTS

Percentage degradation: 10% after 10 days, rising to 53% after 28 days.

CONCLUSIONS

p-Nonylphenol was not readily biodegradable under the test conditions. In this test, p-nonylphenol showed significant biodegradation, but does not meet the OECD 301 B "ready" criteria of 70%. P-nonylphenol is considered inherently biodegradable.

DATA QUALITY

Measurement following OECD 301 B guidelines.

REFERENCES

Draft Nonylphenol SIAR – April 2001 – Summary 2.1

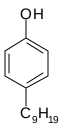
Williams, J.B., Varineau, P.T. 1996. Nonylphenol in biosolids and sludges. SETAC Poster Session P0576, November 20, 1996.

9.3 BIODEGRADATION -ADDITION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



Commercial grade highly branched alkyl chain: 95% p-nonylphenol.

METHOD

Method/guideline followed: OECD 301 F, O₂ Consumption, Manometric Respirometry

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1999.

Contact time: 28 days.

Inoculum: non-adapted, domestic, activated sludge.

Concentration: 31 mg/l (92.4 mg ThOD/l) at 22 °C. No carrier solvents were used.

RESULTS

Percentage degradation: 19% after 10 days, rising to 62% after 28 days.

CONCLUSIONS

p-Nonylphenol was not readily biodegradable under the test conditions. In this test, p-nonylphenol showed significant biodegradation, but does not meet the "ready" criteria due to missing the 10-day window to progress from 10% to 60%. P-nonylphenol is considered inherently biodegradable.

DATA QUALITY

Measurement following OECD 301 F guidelines.

REFERENCES

Draft Nonylphenol SIAR – April 2001 – Summary 2.1

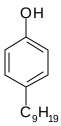
Staples, C.A., et al. 1999. Measuring the biodegradability of nonylphenol ether carboxylates, octylphenol ether carboxylates and nonylphenol," Chemosphere, vol. 38. pp 2029-39, 1999.

9.4 BIODEGRADATION - ADDITION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



METHOD

Method/guideline followed: OECD 301 B - Modified Stürm Test (EEC Directive 79/831 ENV/283/80)

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1996.

Contact time: Hüls 1996(b): 32 days and Hüls 1996(c): 40 days.

Inoculum: (b) non-adapted, domestic, activated sludge (18 x 10⁵ CFU/ml) and (c) adapted, activated sludge (same concentration) (Adaptation: 5 mg/l for 13 days followed by 50 mg/l for 5 weeks). Concentration: (b) and (c): 22.8 mg/l (20 mg carbon/l) with and without an emulsifier.

RESULTS

Percentage degradation: (b) 0% after 32 days, with and without emulsifier, and (c) without the emulsifier: 0% at 40 days, with the emulsifier: 78% at 40 days.

CONCLUSIONS

p-Nonylphenol was not readily biodegradable under the test conditions. The results obtained in the two modified Stürm tests above indicate that NP undergoes biodegradation with adapted microorganisms and could be considered inherently biodegradable. The difference seen in (c) between the results in the presence and in the absence of an emulsifier are difficult to explain.

DATA QUALITY

Measurements following OECD 301 B guidelines with modifications.

REFERENCES

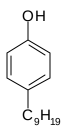
Draft Nonylphenol SIAR – April 2001 – Summary 2.1 Hüls (1996b, 1996c) Determination of the biological degradability of nonylphenol in the modified Stürm test (EEC Directive 79/831 ENV/283/80) Report ST-3 and 3a/84.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance reported as > 95%, confirmed by subsequent gas chromatography.

p-Nonylphenol was provided by Schenectady Chemical Company.

METHOD

Method: No data

Test type: Flow through, unaerated toxicity test

GLP: (Y)

Year study performed: 1990

Species: Cyprinodon variegatus

Analytical monitoring: Yes, details not available on IUCLID

Exposure period: 96 hours

Test conditions: Water quality parameters were within acceptable limits throughout the test.

RESULTS

Nominal concentrations: 0, 0.075, 0.125, 0.19, 0.31 and 0.5 mg/l.

Mean measured concentrations were used for calculations.

LC50 (96hr) 0.31 mg/l

NOEC(96hr) 0.24 mg/l

CONCLUSIONS

The LC50 (96hr) of the test substance is 0.31 mg/l.

DATA QUALITY

GLP study

Information taken from IUCLID database.

REFERENCES

IUCLID Data Sheet

4.1 Acute/Prolonged Toxicity to Fish

Source: ICI Chemicals & Polymers Ltd Runcorn, Cheshire

Huels AG Marl

OTHER

CHRONIC TOXICITY STUDY

IUCLID Data Sheet Chronic Toxicity to Fish

Substance: p-Nonylphenol, purity > 95%

Method: No data

Test type: flow through, unaerated toxicity test

GLP: (Y)

Year: 1991

Species: Pimephales promelas

Exposure period: 33 days

Endpoint: Survival and reproduction rate

Test conditions: Water quality parameters were within acceptable limits throughout the test.

Temperature: 23.4 –26.3°C

Dissolved oxygen: 6.3 - 9.7 mg/l

Results:

Nominal concentrations: 0, 3.0, 6.0, 9.0, 15 and $25 \mu g/l$

Mean measured concentrations were used for all calculations.

NOEC(33days) $7.4 \mu g/l$

LOEC(33days) $14 \mu g/l$

MATC(33days) $10.2 \mu g/l$

Remarks: The most sensitive measured effect was survival of fathead minnows at the conclusion of the test. Fish exposed to the control and the three lowest tested concentration of Nonylphenol (2.8, 4.5 and $7.4 \,\mu g/l$) began to hatch on the third day of exposure, while fish exposed to the highest concentrations

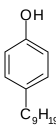
(14 and 23 μ g/l) did not begin to hatch until the fourth day. No statistically significant effects were noted at any test concentration on the number of embryos hatched, the time to first feeding or length and weight of surviving fish. No sublethal effects were noted during the study.

10.2 ACUTE TOXICITY TO FISH -ADDITION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

Method: No data

Test type: Flow-through toxicity test

GLP: (Y)

Year study performed: 1993

Species: Pimephales promelas

Analytical monitoring: Yes, details not available on IUCLID

Exposure period: 96 hours

Test conditions: Have no information – test judged valid in SIAR process.

RESULTS

Nominal concentrations: Have no information – test judged valid in SIAR process.

Mean measured concentrations were used for calculations.

LC50 (96hr) 0.128 mg/l

CONCLUSIONS

The LC50 (96hr) of the test substance is 0.128 mg/l.

DATAQUALITY

Determined to be Valid in SIAR Process

Draft Nonylphenol SIAR – April 2001 – Summary 4.1.1

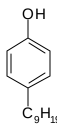
Brooke, L.T. 1993a. Acute and Chronic Toxicity of nonylphenol to ten species of aquatic organisms. USEPA Draft Report, EPA Contract No 68-C1-0034.

10.3 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.13 mg/l

ChV(30 day) 0.017 mg/l

ChV (90 day) 0.005 mg/l Remark: log Kow used 5.92 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.13 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

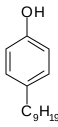
Calculation performed for this HPV submission.

10.4 CHRONIC TOXICITY TO FISH

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

Purity of the test substance reported as > 95%, confirmed by subsequent gas chromatography.

p-Nonylphenol was provided by Schenectady Chemical Company.

METHOD

Method: No data

Test type: Flow through, unaerated toxicity test

GLP: (Y)

Year study performed: 1991

Species: Pimephales promelas

Analytical monitoring: Yes, details not available on IUCLID

Exposure period: 33 days

Endpoint: Survival and Reproduction Rate

Test conditions: Water quality parameters were within acceptable limits throughout the test.

Temperature: 23.4 - 26.3 ℃

Dissolved oxygen: 63 - 9.7 mg/l

RESULTS

Nominal concentrations: 0, 3.0, 6.0, 9.0, 15 and 25 μ g/l.

Mean measured concentrations were used for calculations.

NOEC(33 days) 7.4 μg/l

LOEC(33 days) 14 µg/l

MATC(33 days) 10.2 µg/l

Remarks: The most sensitive measured effect was survival of fathead minnows at the conclusion of the test. Fish exposed to the control and the three lowest tested concentration of Nonylphenol (2.8, 4.5 and 7.4 μ g/l) began to hatch on the third day of exposure, while fish exposed to the highest concentrations (14 and 23 μ g/l) did not begin to hatch until the fourth day. No statistically significant effects were noted at any test concentration on the number of embryos hatched, the time to first feeding or length and weight of surviving fish. No sublethal effects were noted during the study.

CONCLUSIONS

The NOEC (33 days) of the test substance is $7.4 \mu g/l$.

The LOEC (33 days) of the test substance is $14 \mu g/l$.

DATA QUALITY

GLP study

Information taken from IUCLID database.

REFERENCES

IUCLID Data Sheet

4.1 Acute/Prolonged Toxicity to Fish

Source: ICI Chemicals & Polymers Ltd Runcorn, Cheshire

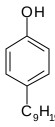
Huels AG Marl

10.5 CHRONIC TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

Purity of the test substance unknown.

METHOD

Method: No data

Test type: Flow through

GLP: Information not provided.

Year study performed: 1993

Species: Pimephales promelas

Analytical monitoring: Yes, details not available on IUCLID

Exposure period: 28 days

Endpoint: Survival

Test conditions: Not provided

RESULTS

NOEC(28 days) = 0.0775 mg/l

LOEC(28 days) = 0.193 mg/l

CONCLUSIONS

The NOEC (28 days) of the test substance is 0.0775 mg/l.

The LOEC (28 days) of the test substance is 0.193 mg/l.

DATA QUALITY

Study deemed valid for NP SIAR - April 2001

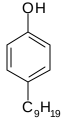
Draft Nonylphenol SIAR – April 2001

Brooke, L.T. (1993b) Accumulation and Lethality for Two Freshwater Fishes (fathead minnow and bluegill) to Nonylphenol. USEPA Draft Report EPA Contract # 68-C1-0034.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance reported as > 95%, confirmed by subsequent gas chromatography.

p-Nonylphenol was provided by Schenectady Chemical Company.

METHOD

Method: No data

Test type: Static, unaerated toxicity test

GLP: (Y)

Year study performed: 1990

Species: Selenastrum capricornutum

Analytical monitoring: Yes, details not available on IUCLID.

Exposure period: 96 hours

Test conditions:

Water quality parameters were within acceptable limits throughout the study. Cell counts were made daily with a hemocytometer.

RESULTS

Nominal concentrations: 0, 0.06, 0.12, 0.25 and 0.5 mg/l.

Mean measured concentrations were used for calculations.

EC50 (96hr) 0.41 mg/l

Remarks: Algae transferred from the test flasks containing the highest test concentration to a flask containing fresh media without nonylphenol, grew from 9,700 to 1,940,000 cells per ml during the 7 days following the conclusion of the test, indicating a lack of algastatic effect.

CONCLUSIONS

The EC50 (96hr) of the test substance is 0.41 mg/l.

DATA QUALITY

GLP study

Information taken from IUCLID database.

REFERENCES

IUCLID Data Sheet

4.3 Toxicity to Aquatic Plants e.g. Algae

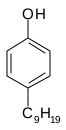
Source: ICI Chemicals & Polymers Ltd Runcorn, Cheshire

Huels AG Marl

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE) - ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance unknown

METHOD

Method: No data

Test type:

GLP:

Year study performed: 1997

Species: Scenedesmus subspicatus (freshwater)

Analytical monitoring: Yes, details not available on IUCLID.

Exposure period: 72 hours

Test conditions: Not provided in summary.

RESULTS

EC50 (72hr) 0.0563 mg/l, based upon change in biomass

CONCLUSIONS

The EC50 (96hr) of the test substance is 0.0563 mg/l.

DATA QUALITY

Study deemed valid in SIAR Process.

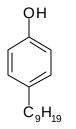
Draft Nonylphenol SIAR – April 2001

Kopf W. (1997). Effect of Endocrine Active Substances in Biotests with Aquatic Organisms. Bavarian State Water Authority, Institute for Water Research, Munich, Oldenburg. (German) (EN 28692/ISO 8692 DIN 384129)

11.3 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE) - ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance: 95%

METHOD

Method: No data

Test type: Static

GLP:

Year study performed: 1990

Species: Skeletonema Costatum (saltwater)

Analytical monitoring: Yes, details not available on IUCLID.

Exposure period: 96 hours

Test conditions: Water Temperature: 21-22 C, pH = 7.9 - 9.6, Salinity: 30%.

RESULTS

EC50 (96hr) 0.027 mg/l, based upon cell growth

CONCLUSIONS

The EC50 (96hr) of the test substance is 0.027 mg/l.

DATA QUALITY

Study deemed valid in SIAR Process.

Draft Nonylphenol SIAR – April 2001

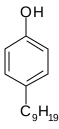
Ward, T.J., Boeri R.L., (1990a) Acute static toxicity of nonylphenol to marine alga (*Skeletonema costatum*). Report prepared for Chemical Manufacturers Association by Resource Analysts. Study No. 8970-CMA.

11.4 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.037 mg/l

ChV (96hr) 0.036 mg/l

Remark: log Kow used 5.92 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.037 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

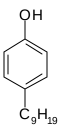
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

Purity of the test substance: information not available.

METHOD

Directive 84/449/EEC, C.2 "Acute Toxicity to Daphnia magna"

Test type: No data

GLP: (Y)

Year study performed: 1992

Species: Daphnia magna (Crustacea)

Analytical monitoring: No

Exposure period: 48 hours

Test conditions: Solvent (acetone)

RESULTS

EC0 (48hr) $< 100 \,\mu g/l$

EC50 (48hr) $140 \mu g/l$

EC100 (48hr) $\geq 400 \,\mu g/l$

CONCLUSIONS

The EC50 (48hr) of the test substance is 140 μ g/l.

DATA QUALITY

GLP study

Information taken from IUCLID database.

Purity of the test substance: information not available.

REFERENCES

IUCLID Data Sheet-

4.2 Acute Toxicity to Aquatic Invertebrates

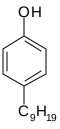
Source: ICI Chemicals & Polymers Ltd, Runcorn, Cheshire

Huels AG Marl

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA) - ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance: information not available.

METHOD

Method: No data

Test type: No data

GLP: No Data

Year study performed: 1993

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 48 hours

Test conditions: No data

RESULTS

EC50 (48 hr) = 0.085 mg/l

CONCLUSIONS

The EC50 (48hr) of the test substance is 0.085 mg/l.

DATA QUALITY

Study deemed valid in SIAR process.

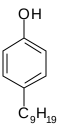
REFERENCES

Draft Nonylphenol SIAR – April 2001. Brooke, L.T. (1993a) Acute and Chronic Toxicity of nonylphenol to ten species of Aquatic Organisms USEPA Draft Report, EPA Contract No. 68-C1-0034.

12.3 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., MYSID) -ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance: information not available.

METHOD

Method: No data

Test type: Flow-Through

GLP: No Data

Year study performed: 1990

Species: Mysidopsis bahia

Analytical monitoring: No data

Exposure period: 96 hours

Test conditions: Temperature of 23.8-25.3 C, Dissolved oxygen of 6.5-7.8 mg/l, Salinity of 20%, and pH range 7.3-8.2.

RESULTS

LC50 (96 hr) = 0.043 mg/l

NOEC (96 hr) = 0.018 mg/l

CONCLUSIONS

The LC50 (96hr) of the test substance is 0.043 mg/l.

DATA QUALITY

Study deemed valid in SIAR process.

Draft Nonylphenol SIAR – April 2001.

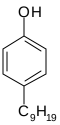
Ward T.J., Boeri, R.L., (1990c) Acute Flow-through Toxicity of nonylphenol to the Mysid (*Mysidopsis bahia*). Report prepared for Chemical manufacturers Association by Resource Analysts. Study No 8974-CMA.

12.4 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.30 mg/l

ChV (21day) 0.014 mg/l

Remark: log Kow used 5.923.28 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.030 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

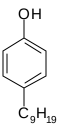
OTHER

Calculation performed for this HPV submission.

12.5 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES (E.G. DAPHNIA) ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance: 91.8% nonylphenol, 86.1% 4-nonylphenol

METHOD

Method: OECD Guideline No. 202, Part 2 "Daphnia sp. Reproduction Test"

Test type: semi-static with endpoint of reproduction rate

GLP: (Y)

Year study performed: 1993

Species: Daphnia magna (Crustacea)

Analytical monitoring: No data

Exposure period: 21 days

Test conditions: Temperature of $20 \pm 1^{\circ}$ C, Hardness 180 ± 20 , pH ± 0.25

0.10 mg/l (0.08 - 0.13 mg/l)

10 replicate vessels, each containing one *Daphnia* were employed for the control, solvent control and each test concentration. The *Daphnia* were fed daily with cultures of algae and yeast.

RESULTS

LC50 (21days)

LC50 values with their 95% confidence limits:

LC50 (1day) 0.30 mg/l

LC50 (2days) 0.19 mg/l (0.13 - 0.25 mg/l)

LC50 (4days) 0.15 mg/l (0.11 - 0.23 mg/l)

LC50 (7days) 0.12 mg/l (0.10 - 0.17 mg/l)

LC50 (14days) 0.12 mg/l (0.09 - 0.16 mg/l)

NOEC (21 day – surviving offspring) 0.024 mg/l

NOEC (21 day - length) 0.039 mg/l

Mean measured concentrations were used for calculations

CONCLUSIONS

The LC50 (21 day - reproduction) of the test substance is 0.10 mg/l.

DATA QUALITY

Study deemed valid – IUCLID and SIAR Process

REFERENCES

IUCLID Data Sheet- Daphnia Reproduction Test

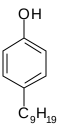
Draft Nonylphenol SIAR – April 2001

Comber M.H.I., Williams, T.D., Stewart, K.M. (1993) The effects of nonylphenol on *Daphnia magna*. Water Research, **27**(2), 273-276.

12.6 CHRONIC TOXICITY TO AQUATIC INVERTEBRATES (E.G., MYSID) - ADDITION

TEST SUBSTANCE

p-Nonylphenol



CAS No. 84852-15-3

IUCLID Identification Number: 84852-15-3

Purity of the test substance: information not available.

METHOD

Method: No data

Test type: Static

GLP: No Data

Year study performed: 1991

Species: Mysidopsis bahia

Analytical monitoring: No data

Exposure period: 28 days

Test conditions: Temperature of 23.3-26.4 C, Dissolved oxygen of 5-8.5 mg/l, Salinity of 20-21%, and pH range 7.5-8.2.

RESULTS

LOEC (length) (28 day) = 0.0067 mg/l

NOEC (length) (28 day) = 0.0039 mg/l

CONCLUSIONS

The LOEC (28 day) of the test substance is 0.0067 mg/l.

DATA QUALITY

Study deemed valid in SIAR process.

Draft Nonylphenol SIAR – April 2001.

Ward T.J., Boeri, R.L., (1991c) Chronic Toxicity of nonylphenol to the Mysid (*Mysidopsis bahia*). Report prepared for Chemical Manufacturers Association by Resource Analysts. Study No 8977-CMA.

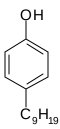
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number 84852-15-3

METHOD

OECD Guidelines No. 401

GLP(Y)

Year study performed: 1981

Species: Rat

RESULTS

LD50 1882 mg/kg

CONCLUSIONS

LD50 1882 mg/kg

DATA QUALITY

To OECD guidelines and GLP compliant.

Information taken from IUCLID database.

REFERENCES

 $IUCLID\ Dataset\ created\ by\ EUROPEAN\ COMMISSION-European\ Chemicals\ Bureau\ created\ 19$ Feb 2000.

OTHER

On IUCLID there is data available for 20 acute oral toxicity tests that have been done in rats over the years. Several of these studies were performed according to OECD guideline 401 and GLP (as example above).

LD50 values were consistently reported to be in the range between 1000 and 2000 mg/kg, with one exception, where a value of 580 mg/kg was reported. However the value of 580 mg/kg relates to an old non-GLP study.

ADDITION: In the Draft Nonylphenol SIAR – April 2001, it is stated that nonylphenol is moderately toxic by the oral route, with LD50 values for the rat in the range of about 1200 to 2400 mg/kg.

REFERENCES - ADDITIONAL

Berol Kemi AB (1982) Nonylphenol acute oral toxicity in rats. Inveresk Research International project no. 230086, report no. 2379.

Hüls AG (1982) Nonylphenol: An Acute Toxicity Study (LD50) in the rat. Hazleton Laboratories Deutschland Project No. 222/8.

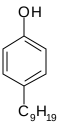
ICI Central Toxicology Laboratory (1984) Nonylphenol (ex-oil Works and Rohm and Haas): Comparison of Acute Oral Toxicities. CTL Report No. CTL/L/708.

13.2 ACUTE TOXICITY SENSORY, SKIN AND EYE IRRITATION

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

SENSORY

REMARK

Acute inhalation studies have not been done to determine an LD50. However assessment of sensory irritation in mice indicates that it is mildly irritating at concentrations of 400ppm.

REFERENCE

IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000

Central Toxicology Laboratory (1995)

Nonylphenol: Assessment of Sensory Irritation potential in mice.

Report Number: CTL/L/6768

Date: 31 August 1996

Study Sponsor: ICI Chemicals & Polymers Limited

SKIN

REMARK

Studies of skin irritation consistently report highly irritating or corrosive properties for nonylphenol. Several of these studies follow current guidelines.

REFERENCE

IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000.

ICI Chemicals & Polymers Limited, Runcorn, Cheshire.

Huels AG Marl

EYE

REMARK

Studies of eye irritation consistently demonstrate the irritation potential of nonylphenol. The only study done to an OECD guideline protocol reports the substance to be irritating.

REFERENCE

IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000.

ICI Chemicals & Polymers Limited Runcorn, Cheshire.

Huels AG Marl

OTHER

ADDITION (taken from Nonylphenol SIAR Draft – April 2001):

Several studies conducted in animals according to OECD 404 are available which indicate liquid nonylphenol can be corrosive to the skin, although its potency might vary according to source and exact composition. In some of these studies, full thickness destruction or skin necrosis were observed within 24 h of a 1 or 4 hour application (Union Carbide, 1992 a,b; Hüls 1986a) whilst in a study sponsored by Enichem (1992), less severe skin reactions were not reversible within an 8 day observation period. In another study (Berol Kemi AB (1987)), skin reactions (grade 2 erythema and grade 1-3 edema) were reversible within 13 days. The liquid is also a severe eye irritant. Two well reported studies using methods equivalent to OECD guideline 405 are available. In one study (Hüls 1986b) maximal scores for conjunctival redness were reported for much of the 21 d observation period and 2 of 3 rabbits tested showed grade 3 or 4 corneal opacities at the end of the observation period. In the second study (ICI, 1979), slightly less severe effects were observed. At the end of the 7 day observation period, eye lesions were still present in two rabbits. Exposure to the saturated vapor (nominally 400 ppm) elicited mild sensory irritation of the respiratory tract in mice, but no reaction was elicited at a nominal concentration of 30 ppm (ICI, 1995).

Sensitisation: The results of several guinea pig maximisation tests (Hüls, 1986c; ICI, 1980, 1979; Gaworski et al., 1979) suggest that nonylphenol does not have significant skin sensitising potential.

REFERENCES - ADDITIONAL

Draft Nonylphenol SIAR - April 2001

Union Carbide (1992 a and b) Nonylphenol RNH: Primary skin irritancy study in the rabbit by Department of Transport (DOT) Procedures. Union Carbide project report 91U0008, and 91U0009.

Hüls AG (1986a) Test of Acute Skin Irritation of Nonylphenol. Hüls Report 0584.

Enichem (1992) Acute Dermal irritation study in Rabbits. Instituto di Ricerche Biomediche Report 910515.

Berol Kemi AB (1987) Irritant effects on rabbit skin of ronylphenol. Huntingdon Research Center Report No. 861361D/BKI 94/SE.

Hüls AG (1986b) Test of Acute Eye Mucus Membrane Irritation of Nonylphenol. Hüls Report 0585.

ICI Central Toxicology Laboratory (1979) Nonylphenol (ex-oil Works and Rohm and Haas): Comparison of acute oral toxicities, skin and eye irritation and skin sensitisation potential. CTL Report No. CTL/T/1278.

ICI Central Toxicology Laboratory (1995) Nonylphenol: Assessment of sensory irritation potential in mice. Zeneca CTL Report No. CTL/L/6768.

Hüls AG (1986c) Skin Sensitisation Effect of Nonylphenol on Guinea Pigs. Hüls Report 0690.

ICI Central Toxicology Laboratory (1980) Nonylphenol samples (ex Rohm and Haas Process): Skin sensitisation studies. CTL Report No. CTL/T/1399.

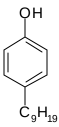
Gaworski C.L., Kimbead E.R., and Doyle, R.L. (1979) Acute toxicity of a number of chemicals of interest to the Air Force. University of California Extension, Wright Patterson Air Force Base, Report ISS AMRL-TR-79-11.

14.1 GENETIC TOXICITY IN VIVO

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

Guideline followed: EEC Directive 79/831, B.12

Type: Micronucleus assay

GLP (N)

Year study performed: 1979

Species/strains: NMRI mice, male and female.

Route of administration: Oral, gavage.

Concentration tested: 500 mg/kg (maximum tolerated dose.)

Monitoring period: 18, 48 and 72 hours.

Nonylphenol was administered once to 10 mice with a dose level of 500 mg/kg.

RESULTS

No mutagenic effects in mouse erythrocytes were observed at any sampling time.

CONCLUSION

The test substance did not demonstrate any mutagenic potential in this in vivo test system.

DATA QUALITY

Study performed to EU guidelines but not GLP.

Information taken from IUCLID database.

Huels-Report 2 Mutagenitaetsuntersuchung von Nonylphenol im Mikrokern-Test", P. Schoeberl, 1988, (unpublished) as cited in IUCLID.

IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000

OTHER

Additional Gene Toxicity - In-Vivo

An *in-vivo* bone marrow micronucleus test in mice, conducted using the intraperitoneal route and at up to a maximally tolerated dose (300 mg/kg), was negative (Hüls, 1999). Re-enforces that nonylphenol is not mutagenic in these test systems.

REFERENCES - ADDITIONAL

Draft Nonylphenol SIAR – APRIL 2001

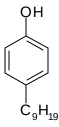
Hüls AG (1999) In vivo mouse micronucleus test. Hüls Report No. MK-99/0255.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID Identification Number: 84852-15-3

METHOD

Guideline followed: Ames test Ames, B.N.: J. McCann, E Yamashi: Mutation Research, 31 347-364, (1975)

Type: Plate incorporation assay

System of testing: Bacterial

GLP (N)

Year study performed: Not known

Species/strains: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537 and TA 1538.

Concentration tested: Dosed up to 5 mg/plate.

Metabolic activation: Dosed with and without metabolic activation.

RESULTS

Negative in both the presence and absence of metabolic activation.

CONCLUSIONS

The test substance was considered to be non-mutagenic in this test system.

DATA QUALITY

Not conducted to GLP.

Data taken from IUCLID.

Huels-Report No. 84/19, Project X41, 1984, (unpublished) as cited in IUCLID

ICI Chemicals and Polymers Limited, Runcorn, Cheshire.

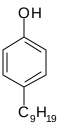
 $IUCLID\ Dataset\ created\ by\ EUROPEAN\ COMMISSION-European\ Chemicals\ Bureau\ created\ 19$ Feb 2000.

15.2 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID ID Number: 84852-15-3

METHOD

Guideline followed: OECD Guideline 476 HGPRT assay

Type: V79 cells from Chinese hamster

System of testing: Non bacterial

GLP (Y)

Year study performed: 1984

Species/Strain: V79 cells from Chinese hamster

Metabolic activation: With and without metabolic activation.

Dose range 0.00016 - 0.01 mg/ml without S9 mix and 0.0004 - 0.1 mg/ml with S9 mix.

RESULTS

No reproducible biologically significant increases in mutant frequency at the HPRT locus were observed when compared to the solvent control.

CONCLUSION

The test substance was tested up to toxic concentrations and is considered to be non-mutagenic in this test system. (This result was confirmed in an independent experiment.)

DATA QUALITY

Study was GLP compliant and conducted according to OECD guidelines.

Information taken from IUCLID.

Draft Nonylphenol SIAR – April 2001.

Huels – Report 688, Final Report: in vitro mammalian cell gene mutation test with nonylphenol, IBR-Project No. 95-86-0446-90, 1990 (unpublished).

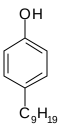
IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000.

16.1 REPEATED DOSE TOXICITY 28 DAY DIET

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID ID Number 84852-15-3

METHOD

OECD Guideline 407

GLP (Y)

Year study performed: 1984

Species: Rats

Strain: Crl:CD(SD)BR

Route of Administration: Diet containing Nonylphenol (purity>=98%)

Duration of test: The animals were treated daily for 28 days.

Doses/concentration levels: nominal dose levels of 0, 25, 100 and 400 mg/kg/day

Four groups of ten animals (5 males and 5 females). These are known as groups 1-4 respectively.

RESULTS

NOAEL: 100 mg/kg/day.

There were no deaths and no clinical signs to suggest any effect of treatment.

Group 4 animals gained less weight and consumed less food than the controls. At week 4 there were changes in the blood chemistry of animals in group 4, (namely decrease in mean glucose level, increase of mean urea and cholesterol level.) There were increases in group 4 kidney, liver and testes weight. Some other slight histopathological changes seen in the males in group 4 at week 4 included hyaline droplet accumulation in the renal proximal tubules, and minor vacuolation in the periportal hepatocytes in the liver. No such treatment related effects were observed in females.

There were no changes in groups 2 and 3 related to treatment.

CONCLUSION

For both males and females the no observable adverse effect level was considered to be $100 \, \text{mg/kg/day}$.

DATA QUALITY

Study conducted according to OECD guidelines and GLP principles.

Information taken from IUCLID.

REFERENCE

Huels-report 774: Nonylphenol: A 28 day oral (dietary) subacute toxicity study in the rat, prepared by Hazleton UK, Report-no. 5917-671/1 (unpublished)

ICI Chemicals and Polymers Limited, Runcorn, Cheshire.

 $IUCLID\ Dataset\ created\ by\ EUROPEAN\ COMMISSION-European\ Chemicals\ Bureau\ created\ 19$ Feb 2000

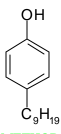
16.2 REPEATED DOSE TOXICITY 90 DAY DIET

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3

IUCLID Identification Number 84852-15-3



METHOD

Guideline followed: U.S. EPA TSCA guidelines

Type: Subchronic Toxicity (90-day) study in rats

GLP (Y)

Year study performed: 1997

Species: Sprague-Dawley rats

Strain: Crl:CD BR

Route of administration: Oral diet

Exposure period: 90 days

Four groups of rats were administered dietary concentrations of 0, 200, 650 or 2000 ppm which corresponded to approximate dietary intakes of 0, 15, 50 or 150 mg/kg/day. There were 25 rats/sex/group in the control and high-dose groups and 15 rats/sex/group in the low and mid-dose groups. 10 of the 25 rats/sex in the control and high-dose groups were designated as recovery animals and were maintained on control diets for 4 weeks after completion of the 90-day exposure period to assess the reversibility of any effects, which might be observed. Estrous cyclicity was monitored during week 8 and sperm count, motility and morphology evaluated at termination to evaluate the possible weak estrogen-like activity that has been reported for the test substance in a number of screening assays.

RESULTS

NOAEL: 50 mg/kg/day (650 ppm in the diet).

No mortality or morbidity occurred.

In-life effects from exposure to the test substance were limited to small decreases in body weight (<10%) and food consumption in the 150 mg/kg/day group. At 90 days, the mean bodyweights for

both sexes at this exposure level were about 7% less than controls. In the satellite group, some recovery of bodyweight and food consumption values were seen after exposure was discontinued. Hematology and opthalmoscopy findings and oestrus cycle patterns spermatogenesis were not affected by treatment. Serum aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels were markedly elevated in two females at 150 mg/kg, which correlates to some histopathological changes reported in the liver.

At necropsy, no treatment related macroscopic findings were reported. Among the males killed at 90 days, there was a dose-related increase in group mean absolute (by 6, 9, and 13%, relative to controls, at 15, 50 and 150 mg/kg/day, respectively) kidney weight. In the recovery group, the bodyweight-related kidney weight among males was also increased, although the effect was less marked. However, this organ weight increase could not be correlated with any clinical chemistry or histopathological change, and consequently, this finding was considered unlikely to be of any toxicological significance, particularly at 15 and 50 mg/kg/day where magnitude of the change was small. Also, ovary weight was slightly decreased in the 150 mg/kg/day group, in comparison with the controls, at 90 days. In contrast, the weight of this organ was slightly increased in the recovery group. Again, this difference could not be correlated with any histopathological change which, together with the inconsistency between the findings for the main and satellite groups, makes the interpretation of this finding uncertain. Body-weight related liver weight was increased at 90 days only in males at 50 and 150 mg/kg/day and females at 150 mg/kg/day, by about 10% compared with controls. This was considered likely to be an adaptive rather than toxicological response.

The only noteworthy microscopic changes were seen in the kidneys and liver. In males at 140 mg/kg/day in both the main and satellite groups, there was a decrease in the occurrence of renal tubular hyaline droplets/ globules in comparison with the control group. The biological significance of this change is uncertain. Also, a lack of a lack of correlation with the findings of the 28 day repeated dose study, in which an actual increase in the incidence of renal hyaline droplets occurred, casts doubt on whether these changes should be considered to be related to treatment. The renal histopathological findings have been reviewed by an independent pathologist (Hard 1998). An increased incidence of deposits of intratubular mineralisation in the P3 (straight) segment of the proximal tubule at the outer stripe of the outer medulla/ inner stripe of the outer medulla junction was seen in males at 150 mg/kg/day; 11 out of 25 from this group were affected, compared with 1 out of 25 control males. Slight or moderate hepatic cell necrosis was seen in three females at 150 mg/kg/day; two of the affected females also had raised serum ALT and AST. This provides evidence that the liver may be a target organ for nonylphenol toxicity, although this evidence is weak in view of the mild nature of response and small number of animals affected.

Postmortem measurements at Week 14 indicated a dose-related kidney weight increase in males and a decrease in renal hyaline globules/droplets in males from the high-dose group. The kidney weights showed complete recovery following a 4-week recovery period. The changes were of small magnitude and there were no corresponding clinical or histopathological changes, and thus these findings were not regarded as toxicologically significant.

No other effects attributable to the test substance were observed.

No changes were observed for estrous cycling, sperm evaluations or effects on endocrine organs.

CONCLUSION

Based on the minor findings for the 150 mg/kg/day dose group, the NOAEL was considered to be 650 ppm in the diet = 50 mg/kg/day. At 150 mg/kg/day, there were reductions in bodyweight gain, food consumption and food utilisation together with evidence of morphological changes in the liver and possibly kidneys.

DATA QUALITY

Conducted according to EPA guidelines and GLP.

REFERENCE

Draft Nonylphenol SIAR – April 2001.

Cunny, H.C., Mayes B.A., Rosica, K.A., Trutter, J.A., VanMiller, J.P. (1997) Subchronic toxicity study with para-nonylphenol in rats. Reg. Toxicol. Pharmacol. 26, No. 2, pp 172-178.

Chemical Manufacturers Association (1997a) 90-day dietary study in rats administered paranonylphenol. Corning Hazelton study CHV 2603-105.

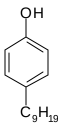
Hard G.C. (1998) Expert report on renal histopathologic changes in rat dietary studies with nonylphenol. Report prepared for the Alkylphenol and Ethoxylates Research Council, Washington D.C., USA.

17.1 REPRODUCTIVE TOXICITY

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID ID Number 84852-15-3

METHOD

Guideline followed: National Institute of Environmental Health Sciences (NIEHS)

Type: Three generation Reproduction study

GLP (Y)

Year study performed: 1997

Species: Sprague-Dawley rats of the Crl:CD BR strain

Route of administration: Oral diet

Doses levels: 0, 200, 650 and 2000 ppm, corresponding to approximately 0,15, 50 and 140 mg/kg/day.

Number of animals and sex: 30 male and 30 female per group.

Age of animals before initiation of dosing: 7 weeks

Frequency of treatment: Continuous from study day 1 until necropsy/termination

Premating exposure for F₀ generation males and females: 6 weeks

F₁ and F₂Generations

The dams reared their litters until weaning on Postnatal Day (PND) 21.

At birth pups were sexed, counted and weighed and anogenital distance was measured.

30 pups/sex/group selected for mating trial and 20 pups/sex/group for PND 21 necropsy.

The day of testicular descent, vaginal opening and preputial separation determined.

 F_1 and F_2 animals received diet containing the doses after weaning and at the same dose as their parents.

Generations raised until mating at sexual maturity (PND 86 ± 10).

Vaginal smears obtained for three weeks prior to cohabiting.

Cohabiting for up to 14 days, pregnancy confirmed by testing for vaginal sperm.

On completion of lactation phase adults were weighed and necropsied, organ weights obtained, sperm analysis performed and tissues fixed and saved. Tissues preserved were liver, kidneys, spleen, stomach, duodenum, jejunem, ileum, ventral prostate, seminal vesicles with coagulating glands, gros s lesions, vagina/cervix, uterus and ovaries or testis and epididymis.

Animals observed twice daily for mortality and signs of toxicity. Body weights, physical examinations and feed consumption obtained every two weeks after weaning.

F₃Generation

As above until study terminated upon sexual development observations of F₃generation (PND 55-58).

Statistical methods:

ANOVA and Dunnett's test used to measure tests for homogeneity of variances.

Hypotheses were tested using nonparametric multiple comparisons procedure of Dunn (1964) or Shirley (1977), as modified by Williams (1986). Also used were Jonckheers's test (1954) and Wilcoxon's test.

Cochran-Armitage test used to test for a dose-related trend, and pairwise comparisons were performed using a chi-square test (Conover, 1971).

Equality of pup weight was evaluated using a parametric analysis of covariance (Nester and Wasserman, 1974).

RESULTS

The overall average estimated dosage during non-reproductive phases for the 200, 650 and 2000 ppm animals varied from 13-19 mg/kg/day, 43-64 mg/kg/day and 131-199 mg/kg/day respectively. The overall estimated dosage during lactation varied from 27-30 mg/kg/day, 93-98 mg/kg/day and 274-322 mg/kg/day respectively.

Maximum tolerated dose was 2000 ppm based on decreased body weight gain and the presence of renal lesions.

Adult body weights of animals dosed at 650 ppm, were reduced in both sexes, at some but not all timepoints. Adult body weights of animals dosed at 2000 ppm animals were reduced in all generations and in both sexes, except F_0 males.

 F_0

Although a few clinical signs were observed the incidence of these was low and no dose-related differences were observed.

Fertility was not affected.

No dose related macroscopic or microscopic lesions were seen at in the adults at necropsy.

 \mathbf{F}_1

PND 21 Necropsy-

Increases in weight of vagina/cervix/uterus in the 650 and 2000 ppm group. Gross lesions noted in one 2000 ppm male and one 2000 ppm male.

Mating trial

Day of vaginal opening accelerated in 650 and 2000 ppm F_1 females. Preputial separation delayed in 2000 ppm F_1 males. No differences in day of testicular descent.

Treatment related decreases in body weight in both male and females were seen at 650 and 2000 ppm.

No dose-related differences in clinical signs were observed: incidences were low to moderate (less than 20% per group).

Continued exposure did not affect the fertility of the F_1 animals, litter size, gestation length, proportion of live born pups, sex ratio, live F_2 pup weight adjusted for litter size, pup survival, anogenital distance, or live pup weights.

Increased length of the estrous cycle and decrease in number of cycles was seen in 2000-ppm F_1 females.

Morbidity was seen in only one 650 ppm F₁ female and this was due to mastitis.

Necropsy Results

Mean terminal body weights and mean absolute weights for the ventral prostrate and right testis of the 2000 ppm F_1 males were decreased. Absolute kidney weights and relative kidney weights were also increased for this group. The mean terminal body weights of the 650 and 2000 ppm F_1 females were also decreased, as were the mean absolute weights of the liver and ovaries. The relative kidney weight was increased for the 2000-ppm F_1 females.

No dose related changes were seen regarding sperm analysis.

No treatment related lesions were noted.

Microscopic evaluation of the kidneys revealed treatment related lesions in the F₁ males and females.

 F_2

PND 21 Necropsy

Incidence of gross lesions was very low and no dose response was observed. Relative kidney weights increased by 9% in 2000 ppm group F_2 females.

Mating Trial

The day of vaginal opening was accelerated in 650 and 2000 ppm females and testicular descent was accelerated. No differences were observed in the day of preputial separation.

Treatment related decreases in body weight were observed in the 650 and 2000 ppm males and in the 2000 ppm females.

Mean feed consumption values were decreased in all F₂ males at some time during the study, and some females.

No dose related clinical signs were observed. Incidences were low.

Continued exposure did not affect the fertility of the F_2 animals, litter size, gestation length, proportion of live born pups, sex ratio, live F_3 pup weight adjusted for litter size, pup survival, anogenital distance.

An increase in cycle length was seen in the 2000 ppm F₂ females.

Decreased epididymal sperm density and testicular spermatid head counts were seen in the F₂ generation.

Adult Necropsy results

Some changes in organ weights seen. Ovarian weights were decreased at 650 ppm in F_2 generation and at 2000 ppm in the F_1 F_2 and F_3 generations.

No differences seen in sperm analysis parameters.

Microscopic evaluation of kidneys revealed treatment-related lesions in males and females.

 F_3

PND 21 necropsy

 F_3 pup terminal body weights were decreased at 650 in males and females. Changes seen in epididymis (relative weight of the right cauda epididymis increased by 24%) and spleen weights (decreased by 24%).

The day of vaginal opening was accelerated in the 650 and 2000 ppm F_3 females. The day of testicular descent was delayed in males. No differences were observed in the day of preputial separation.

F₃ Terminal necropsy

Mean body weights of the 2000 ppm F_3 males decreased. For the 650 and 2000 ppm males the right epididymis were decreased. Testis to terminal body weight ratio increased in 2000 ppm.

Mean body weights of the 650 and 2000 ppm F₃ females decreased. Also decreased were absolute weights of liver, kidneys and vagina/cervix/uterus in the 650 and 2000 ppm females.

No dose related lesions recorded.

Microscopic evaluation of kidneys revealed dose related lesions in males and females.

RESULTS AS SUMMARIZED IN THE DRAFT NONYLPHENOL SIAR - APRIL 2001:

Evidence of general toxicity was seen in adults of all generations, although there were no treatment-related clinical signs, mortalities or adverse effects on food consumption. At 160 mg/kg/day, bodyweight gain was reduced in comparison with controls in adults across all generations, with the terminal bodyweights being about 10% lower than in controls. Similar reductions in bodyweight gain were also seen at 50 mg/kg/day in F1 females, F2 males and F3 females. Relative kidney weights were increased at 50 and/or 160 mg/kg/day in adult males of the F0, F1, and F2 generations and also at 160 mg/kg/day in F1 adult females. Histopathological examination revealed an increase, although often without a convincing dose-response relationship, in the incidence of renal tubular degeneration and/or dilation in adult males from all generations and all nonylphenol treated groups; similar findings were reported for adult females at 160 mg/kg/day in the F1, F2, and F3 generations and at 15 and 50 mg/kg/day in the F3 generation. It is difficult to decide for certain whether or not this increased incidence of renal tubular degeneration and/or dilation is related to treatment because these changes were not seen to the same extent in the 90-dasy study, which was conducted using the same strain of rats, and because a dose-dependant trend was not apparent in all generations/ sexes. The lack of

concordance between the studies cannot be explained on the basis of a slightly longer exposure period in the multigeneration study because kidney effects were seen in the F3 generation which were exposed for only 8 weeks, nor on the basis of *in utero* and neonatal exposure because the effect also occurred in the F0 generation. Giving special emphasis to the fact that the increased incidence occurred consistently across all four generations in the multigeneration study, it is considered that this cannot be dismissed as background variation.

The renal histopathological findings have been reviewed by a pathologist not involved in the original investigation (Hard, 1988). The presence of renal lesions in all nonylphenol exposed groups was confirmed, as was the lack of consistent dose-dependent trend in all generations. The predominant renal lesions were described as tubular mineralisation at the OSOM/ISOM junction, cystic tubules surrounded by fibrosis, or granular cast formation at the OSOM/ISOM junction.

CONCLUSION

Reproductive changes seen in both males and females at ≥ 650 ppm, although there was no consistent effect on fertility in the F_0 , F_1 or F_2 mating trials.

Estrous cycle was lengthened (2000 ppm) and a clear treatment-related change in the day of vaginal opening was seen in all three generations. The acceleration in vaginal opening was taken as an indication of the estrogenicity of the test substance, and was considered not necessarily adverse in itself.

No changes were noted in the remaining reproductive/developmental parameters including pregnancy index, mating index, proportion of pups born alive, sex ratio, pup weights, anogenital distance and survival.

No clear treatment-related changes were noted in testicular descent, preputial separation or sex ratio.

Decreased epididymal sperm density and testicular spermatid head counts were seen in the F_2 generation. These changes were not linked to alterations in fertility.

NOAEL – Systemic and reproductive toxicity = 200 ppm = approximately 15 mg/kg/day.

In multigeneration studies in the rat involving dietary exposure for up to 20 weeks, a LOAEL for repeated dose of 15 mg/kg/day was identified, based on histopathological changes in the kidneys (tubular degeneration or dilation), although such changes were not apparent at this dose level in a 90-day dietary rat study. At higher dose levels the liver may also be a target organ; minor histopathological changes in the liver (Vacuolation in the periportal hepatocytes or occasional individual cell necrosis) were seen at doses of 140 mg/kg/day and above in some studies. The oral toxicity of nonylphenol appears to be enhanced when dosed by gavage, with mortalities being reported at dose levels of 100 mg/kg/day and above.

DATA QUALITY

Study performed using NIEHS guidelines and to GLP.

REFERENCES

Final report on the reproductive toxicity of nonylphenol administered by gavage to Sprague-Dawley rats.

National Toxicology Program National institute of Environmental Health Sciences P.O. Box 12233 Research Triangle Park, NC 27709

OTHER

Please see summary 16.2.

Toxicokinetic Studies

Weak estrogenic activity has been reported for p-alkylphenols with relative potencies several orders of magnitude lower than that observed for the physiological hormone 17ß-estradiol. Concerns have been raised concerning bioaccumulation in the blood of humans exposed to trace levels of alkylphenols. Toxicokinetic studies with p-Nonylphenol as a model have been conducted in rats to address these concerns. The results suggest that the test substance does not bioaccumulate in rats receiving low oral doses and only if the detoxification pathways in the liver are saturated by excessive doses does bioaccumulation occur.

Estrogenic Activity

Several assays have been performed using p-nonylphenol in a number of *in vitro* and *in vivo* test systems to help identify estrogen-like activity.

The main conclusions from these studies are that increases in uterine weight were only seen at very high doses compared to the positive control.

SUMMARY FROM DRAFT NONYLPHENOL SIAR – APRIL 2001

No human data are available. Nonylphenol has been shown to have estrogenic activity in a number of in vitro (Routledge and Sumpter, 1996; Soto et al 1991; White et al 1994) and in vivo assays (ICI, 1996; CMA 1997b; Lee and Lee, 1996). The potency of this estrogenic activity In these assays ranged from 3 to 6 orders of magnitude less than that of estradiol. The effects of nonvlphenol on fertility and reproductive performance have been investigated in a good quality oral multigeneration study in the rat at doses 0, 15, 50 and 160 mg/kg/day (NTP, 1997). This study provided evidence that nonylphenol exposure over several generations can cause minor perturbations in the reproductive system of offspring, namely slight changes in the estrus cycle length (15% at 160 mg/kg/day) and possible also in ovarian weight and sperm/spermatid count, although functional changes in reproduction were not induced at the dose levels tested. The NOAEL for these changes was 15 mg/kg/day. The observed perturbations in offspring are compatible with the predictable or hypothesised effects of exogenous estrogenic activity. Evidence of testicular toxicity, seen as seminiferous tubule vacuolation, cell necrosis and a reduction in tubule diameter, was reported at exposure levels which also cause mortality in a repeat dose gavage study in rats (deJager, 1999a). The LOAEL for testicular toxicity was 100 mg/kg/day. This indicates that the toxicity of nonylphenol appears to be enhanced by gavage administration in comparison to dietary administration, presumably because higher peak blood concentrations on nonylphenol are achieved by gavage. No evidence that nonylphenol is a developmental toxicant was seen in a standard oral development toxicity study in rats dosed at 0, 75. 150, 300 and 600 mg/kg/day on gestation days 6.15 (Initiative Umweltrelevante Alstoffe, 1992). Maternal toxicity was manifested as macroscopic changes to the liver and spleen at = 150 mg/kg and mortalities at 300 mg/kg. In contrast, in a gavage study, involving in utero, lactational and direct postweaning exposure at daily doses of 0, 100, 250, and 400 mg/kg, there was evidence of a reduction in sperm count (36%) at 250 mg/kg/day, although it is not possible to state whether this is a developmental effect or as a result of direct exposure after weaning (deJager et al, 1999b). No offspring were born from mothers receiving 400 mg/kg/day; it is not clear from the report if this was because of maternal deaths or embryonic/ fetal resorption. In an intraperitoneal study designed to investigate the effects of nonylphenol on male reproductive tract development of neonatal rats, evidence of impaired development was observed (Lee, 1998). However, this study was difficult to interpret, such that those results carry little weight in the overall assessment of the available data. Overall, the observations of estrogenic activity in the in vitro and in vivo assays, minor perturbations in the reproductive system of offspring in the multigeneration study, and testicular changes in gavage studies collectively raise concerns for reproductive toxicity, possibly mediated through action on the estrogen receptor.

ADDITIONAL REFERENCES

Draft Nonylphenol SIAR – April 2001:
Chemical Manufacturer's Association (1997b) Uterine weight assay of p-nonylphenol and p-octylphenol ethoxylate-5 (OPE-5) administered orally to ovariectomized Sprague Dawley rats. MB Research Labs Project No. MB-96-4960.07
Initiative Umweltrelevante Altstoffe (1992) Teratogenicity study in Wistar rats treated orally with nonylphenol. IBR Project No. 20-04-0502/00-91.
Lee, P.C. (1998) Disruption of male reproductive tract development by administration of xenoestrogen, nonylphenol, to newborn rats. Endocrine 9, 105-111.
Lee, P.C., and Lee W. (1996) In vivo estrogenic action of nonylphenol in immature female rats. Bull Environ. Contam. Toxicol. 57, 341-348.
241-248.

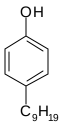
White R., Jobling, S., Hoare, S.A., Sumpter, J.P., Parker M.G. (1994). Environmentally persistent alkylphenolic compounds are estrogenic. Endocrinology, 135(1), 175-182.

18.1 DEVELOPMENTAL TOXICITY/TERATOGENICITY

TEST SUBSTANCE

p-Nonylphenol

CAS No. 84852-15-3



IUCLID ID Number 84852-15-3

METHOD

Guideline followed: EEC directive 87/302, Part B, p.24

Type: "Teratogenicity test-rodent and non-rodent"

GLP (Y)

Year study performed: 1981

Species: Rat

Strain: Wistar

Route of administration: Oral gavage

Doses/Concentrations: 75, 150 and 300 mg/kg/day

Sex: Female

Exposure period: 6-15 day of gestation

Frequency of treatment: Daily

Control group: Yes

Duration of test: 20 days

RESULTS

NOAEL maternal toxicity: 75 mg/kg/day

NOAEL developmental toxicity: 300 mg/kg/day

Treatment of pregnant females from Day 6 to Day 15 of gestation at a dose level of 75 mg/kg was without any general toxicological effect. At a dose level of 150 mg/kg/day 3 of 21 females showed pale and irregularly shaped kidneys, reddening of the renal pelvis and small spleens. 300 mg/kg/day caused clear maternal toxic effects, i.e. increased mortality, reduced body weight gain and food

Page 292 of 317

consumption, and kidney and spleen effects. Caesarean sections were carried out on day 20 of gestation. The findings did not disclose any biologically significant differences between groups in the mean number and presentation of the fetuses, the left or right intra-uterine distribution, the sex ratio, fetal and placental weights, the number of runts and dead fetuses, resorptions, implantation and corpora lutea as indices. Fetal examination did not disclose any treatment related malformations or abnormalities. With regard to the embryo-fetal development a no observable adverse effect level of 300 mg/kg/day was found.

CONCLUSION

Test substance did not show teratogenic effects in rats.

DATA QUALITY

The study was performed according to current EU guidelines and to GLP.

Information taken from IUCLID database

REFERENCE

IBR Forschungs GmbH, D-3030 Waldsrobe: IBR-Project-No: 20-04-0502/00-91 (1992)

Sponsor: INITATIVE UMWELTRELEVANTE ALTSTOFFE E.V. Kennedyalle 93, W-6000 Frankfurt 70 as cited in IUCLID database

IUCLID Dataset created by EUROPEAN COMMISSION – European Chemicals Bureau created 19 Feb 2000

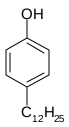
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Melting point: 101.68°C.

CONCLUSIONS

The test substance has a calculated melting point of 101.68°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30 $\,$

OTHER

Weighted value used.

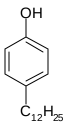
This substance is a mixture of isomers. A representative structure has been used for this calculation.

2.1 BOILING POINT

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: not stated.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Boiling point: 308°C.

CONCLUSIONS

The test substance has a boiling point of 308° C (atmospheric pressure assumed).

DATA QUALITY

Not a GLP study.

Information taken from a safety datasheet.

Method/purity/decomposition: information not available.

REFERENCES

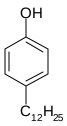
Schenectady International Inc. Material Safety Datasheet (1/19/99).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: $2.09 \times 10^{-5} \text{ mm Hg } @ 25^{\circ}\text{C} (0.0028 \text{ Pa}).$

CONCLUSIONS

The test substance has a calculated vapor pressure of 2.09 x 10⁻⁵mm Hg @ 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30.

OTHER

Modified Grain method.

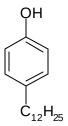
This substance is a mixture of isomers. A representative structure has been used for this calculation.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 7.17.

CONCLUSIONS

The test substance has a calculated log Kow of 7.17.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

This substance is a mixture of isomers. A representative structure has been used for this calculation.

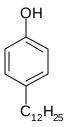
KOWWIN reported an experimental log Kow of 7.91 in its database but this was for a *p*-dodecylphenol with a CAS no. of 104-43-8 (Itokawa et al; 1989).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: calculation using a log Kow value of 7.17.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: $0.05811 \text{ mg/l} @ 25^{\circ}\text{C}$.

CONCLUSIONS

The test substance has a calculated water solubility of 0.05811 mg/l @ 25°C.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

This substance is a mixture of isomers. A representative structure has been used for this calculation.

The log Kow value (7.17) calculated using the programme KOWWIN (see 4.1) has been used for the calculation of water solubility.

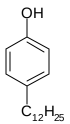
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $48.4 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.65 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.65 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

OTHER - ADDITIONAL

IUCLID Reference on CAS # 121158-58-5, dodecylphenol, branched: half-life = 5.2 hours (indirect photolysis, OH radicals) Reference listed as IUCLID, confidential in "An Initial Review of Potential Risks from Alkylphenols other than Nonylphenol," produced under contract for the UK Environment Agency by WRc-NSF, UK.

7.1 STABILITY IN WATER

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

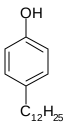
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 262
Data temperature (°C): 25
Water solubility (mg/l): 0.05811
Vapor pressure (Pa): 0.0028
Log Kow: 7.17
Melting point (°C): 102

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	0.019
Soil	97.7
Water	0.0075
Suspended sediment	0.068
Fish	0.0055
Sediment	2.17

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

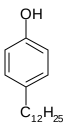
This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method/guideline followed: OECD 301 B, CO₂ Evolution, Modified Stürm Test

Test type: aerobic, measuring CO₂ evolution compared to theoretical amount for complete oxidation.

GLP (Y/N): no.

Year study performed: 1992.

Contact time: 28 days.

Inoculum: non-adapted, domestic, activated sludge from municipal sewage plant; 3300 colony

forming units per ml.

Concentration: 10 mg/l and 20 mg/l

Conditions: Temperature of 21-23 C, no emulsifier

Control: Sodium Benzoate

RESULTS

Percentage degradation: 25% at 10 mg/l after 28 days

6% at 20 mg/l after 28 days

Control Sodium Benzoate achieved 95% degradation within 28 days, reaching threshold for ready biodegradation within 14 days. This indicates inoculum had sufficient biological activity.

CONCLUSIONS

Dodecylphenol, without emulsifier, achieved a degradation level of 25% at 10 mg/l and 6% at 20 mg/l within a period of 28 days. This result indicates that dodecylphenol is not readily biodegradable under the conditions of this test and consequently, dodecylphenol is not considered readily biodegradable. This material is considered inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

Measurement following OECD 301 B guidelines.

REFERENCES

Hüls (1992 a) as cited in the Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

9.2 BIODEGRADATION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in weeks

Ultimate biodegradation in weeks/months.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

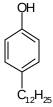
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.025 mg/l

ChV (30 day) 0.003 mg/l

ChV (90 day) 0.00153 mg/l

Remarks: log Kow used 7.17 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.025 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

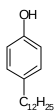
Calc ulation performed for this HPV submission.

10.2 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

No data

GLP: (no data)

Year study performed: unknown

Species: Leuciscus idus (Golden orfe)

Exposure period: 96 hours

RESULTS

NOEC (96hr) 0.5 mg/l

No effects of mortality evident at concentrations up to approximately 0.5 mg/l.

CONCLUSIONS

Experimental NOEC (96hr) for the test substance was found to be 0.5 mg/l.

DATA QUALITY

Data assessed as being valid in the draft RAR. Based on the technical reliability of this study, this result was used in the RAR as an LC50 instead of the using the less reliable, lower salmon value.

IUCLID

REFERENCES

IUCLID Reference on CAS # 121158-58-5, dodecylphenol, branched: Reference listed in "An Initial Review of Potential Risks from Alkylphenols other than Nonylphenol," produced under contract for the UK Environment Agency by WRc-NSF, UK.

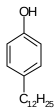
Hüls (1992 a) as cited in Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

10.3 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Method: static

GLP: (no data)

Year study performed: McLeese et al. 1982

Species: Salmo salar (Atlantic Salmon)

Exposure period: 96 hours

RESULTS

LC50 (96hr) 0.14 mg/l

CONCLUSIONS

Experimental LC50 (96hr) for the test substance was found to be 0.14 mg/l.

DATA QUALITY

The data from this study must be used with care as it does not meet the quality criteria since there were insufficient test organisms used, the exposure concentration decreased exponentially during the test and the study was not performed according to GLP. (Draft RAR)

IUCLID

REFERENCES

IUCLID Reference on CAS # 121158-58-5, dodecylphenol, branched: Reference listed as McLeese et al (1982) in "An Initial Review of Potential Risks from Alkylphenols other than Nonylphenol," produced under contract for the UK Environment Agency by WRc-NSF, UK.

Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

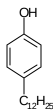
11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE) - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5

Purity: 99.9%



METHOD

Type: Static, measuring biomass and growth rate

GLP: (N)

Year study performed: 2000

Species: Scenedesmus subspicatus

Exposure period: 72 hours

Test Conditions: 22 ± 1 , pH = 9.6-9.7

RESULTS

EC10 (72hr) (growth) 0.53 mg/l

EC50 (72 hr) (growth) 0.77 mg/l

EC10 (72 hr) (biomass) 0.77 mg/l

EC50 (72 hr) (biomass) 0.77 mg/l

72 hour NOEC 0.44 mg/l

CONCLUSIONS

EC50 (72 hr) based on biomass was found to be 0.77 mg/l.

72 hour NOEC was found to be 0.44 mg/l.

DATA QUALITY

Deemed valid through the RAR Process.

REFERENCES

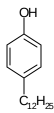
Hüls AG (1992d) as cited in the Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.003 mg/l

ChV (96hr) 0.007 mg/l

Remark: log Kow used 7.17 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.003 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

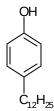
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA) - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Static

GLP: (N)

Year study performed: 1992

Species: Daphnia Magna (Daphnid)

Exposure period: 24 hours, 48 hours

Test Conditions: Temperature of 20 ± 1 C, Dissolved oxygen of >60 mg/l and pH of 7.9.

RESULTS

EC50 (24hr) 0.11 mg/l

EC50 (48 hr) 0.093 mg/l

EC10 (48 hr) 0.056

CONCLUSIONS

Measured EC50 (48hr) for the test substance was found to be 0.11 mg/l.

DATA QUALITY

Deemed valid in RAR Process

REFERENCES

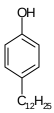
Hüls AG (1992c) as cited in the Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., Sand Shrimp) - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Semi-Static

GLP: (N)

Year study performed: 1981

Species: Crangon septemspinosa (Sand Shrimp)

Exposure period: 96 hours

Test Conditions: Temperature of 10 C.

RESULTS

LC50 (96hr) 0.15 mg/l

CONCLUSIONS

Measured LC50 (96hr) for the test substance was found to be 0.15 mg/l.

DATA QUALITY

Deemed "Use with care" in RAR Process: Although the only data for a marine organism, this study should be used with care, as it does not meet the criteria for a fully valid study due to small size and insufficient number of test organisms.

REFERENCES

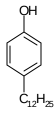
McLeese et al. (1981) as cited in the Targeted Environment Risk Assessment Report for Dodecylphenol, Mixed Isomers (branched) – draft – June 2001. Prepared by the Environment Agency in the UK by WRc-NSF Ltd.

12.3 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.11 mg/l

ChV (21day) 0.003 mg/l

Remarks: log Kow used 7.17 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.11 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

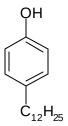
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL - ADDITION

TEST SUBSTANCE

p-Dodecylphenol

CAS No. 210555-94-5



METHOD

Methodology Unknown – IUCLID Confidential - unpublished

GLP(no data)

Year study performed: unknown

Species: Unknown

RESULTS

LD50 2100 mg/kg bw

CONCLUSIONS

LD50 2100 mg/kg bw

DATA QUALITY

Information taken from IUCLID database – IUCLID Confidential, Unpublished.

REFERENCES

IUCLID Reference on CAS # 121158-58-5, dodecylphenol, branched: Reference listed as IUCLID Confidential, unpublished in "An Initial Review of Potential Risks from Alkylphenols other than Nonylphenol," produced under contract for the UK Environment Agency by WRc-NSF, UK.

HEALTH ELEMENTS

p-Dodecylphenol

CAS No. 210555-94-5

No additional test reports found

Alkylphenols Category SECTION FOUR Di-, and Tri-Substituted Mixed Alkylphenols **Chemical Right-to-Know Initiative HPV Challenge Program** Submitted to US EPA by: Schenectady International 2750 Balltown Road New York 12309 USA

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: 1994.

RESULTS

Melting point: 60-62°C.

CONCLUSIONS

The test substance has a melting point range of 60-62℃.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

BASF AG, Sicherheitsdatensblatt Trimethylphenol fest (01.02.1994) as cited in IUCLID database.

1.2 MELTING POINT

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): No.

Year study performed: 1998.

RESULTS

Melting point: 64.9°C.

CONCLUSIONS

The test substance has a melting point of 64.9°C.

DATA QUALITY

Not a GLP study.

Information taken from a technical datasheet.

Purity of the test substance given as typically 99.3% minimum.

Decomposition: no information available.

REFERENCES

Schenectady International Inc. Technical Datasheet (06.11.98).

2.1 BOILING POINT

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1994.

RESULTS

Boiling point: 215°C.

Pressure: assumed to be atmospheric.

CONCLUSIONS

The test substance has a boiling point of 215° C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

BASF AG, Sicherheitsdatensblatt Trimethylphenol fest (01.02.1994) as cited in IUCLID database.

2.2 BOILING POINT

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1998.

RESULTS

Boiling point: 222°C @ 760 mm Hg.

CONCLUSIONS

The test substance has a boiling point of 222° C @ 760 mm Hg.

DATA QUALITY

Not a GLP study.

Information taken from a technical datasheet.

Purity of the test substance given as typically 99.3% minimum.

Decomposition: no information available.

REFERENCES

Schenectady International Technical Datasheet (06/11/98).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1994.

RESULTS

Vapor pressure: < 10 Pa @ 20℃.

CONCLUSIONS

The test substance has a vapor pressure of : <10 Pa @ 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

BASF AG, Sicherheitsdatensblatt Trimethylphenol fest (01.02.1994) as cited in IUCLID database.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: OECD Guideline 107 "Partition Coefficient (n-octanol/water), Flask-shaking Method.

GLP (Y/N): not known.

Year study performed: not known.

RESULTS

Log Kow: 2.72 at 25°.

CONCLUSIONS

The test substance has a log Kow of 2.72 at 25 °C.

DATA QUALITY

Method follows OECD guideline. Temperature but not test substance purity given.

Information taken from IUCLID database.

REFERENCES

BASF AG, Analytisches Labor, unveroeffentlichte Unterschung (J. Nr. 129299/03 vom 01.07.88) as cited in IUCLID Database.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Water solubility: 1.42 g/l at 25°C.

CONCLUSIONS

The solubility of the test substance in water is 1.42 g/l at 25°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance: information not available in database.

REFERENCES

BASF AG, Analtyisches Labor, unveroeffentlichte Untersuchung (J. Nr. 107545/01 vom 07.06.89) as cited in IUCLID Database.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: 131×10^{-12} c m³/molecule-sec.

Half-life: 58.7 minutes.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 58.7 minutes.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

The IUCLID database contains a review of a paper (Faust, B.C., Hoigne, J., Environ. Sci. Tech.21, 957-962 (1987)) on indirect photolysis of the test substance in water but gives no details as to method or results. It does however contain the following summary:

"In natural sunlit waters, 2,3,6-trimethyl phenol is expected to rapidly photooxidize. Photooxidation half-lifes of the 2,4,6,-trimethyl phenol isomer in mid-latitude surface waters during mid-summer are typically on the order of 4-11 hours, and half-lifes for the 2,3,6-trimethyl phenol isomer are expected to be on the same order of magnitude."

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

OTHER - ADDITIONAL

In another test cited by IUCLID, the half-life of this material was determined to be 5 hours, based on photolysis and OH radicals.

IUCLID (non-confidential) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

OTHER - ADDITIONAL

"In natural sunlit waters, 2,3,6-trimethyl phenol is expected to rapidly photooxidize. Photooxidation half-lifes of the 2,4,6,-trimethylphenol isomer in mid-latitude surface waters during mid-summer are typically on the order of 4-11 hours, and half-lifes for the 2,3,6-trimethyl phenol isomer are expected to be on the same order of magnitude."

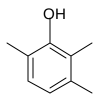
IUCLID (non-confidential) as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



COMMENT

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 136
Data temperature (°C): 25
Water solubility (mg/l): 1420
Vapor pressure (Pa): 10
Log Kow: 2.72
Melting point (C): 64.9

Environmental conditions: defaults used.

RESULTS

Environmental compartment	percentage of test substance
Air	11.6
Soil	27.9
Water	59.9
Suspended sediment	0.019
Fish	0.0016
Sediment	0.62

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

OTHER

9.1 BIODEGRADATION

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



METHOD

Method/guideline followed: Zahn-Wellens Test.

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1981.

Contact time: 14 days.

Inoculum: activated sludge.

Concentration: 1260 mg/l.

RESULTS

Percentage degradation: 6% after 3 hours, 64% after 7 days and 98% after 14 days.

CONCLUSIONS

2,3,6-Trimethylphenol was inherently biodegradable under the conditions of the study.

DATA QUALITY

Not a GLP study.

REFERENCES

BASF AG, Labor fuer Abbau und Analytik; unveroeffentlichte Mitteilung vom 11.12.90 (75076/1981) as cited in IUCLID database.

OTHER - ADDITIONAL

In another biodegradability test; aerobic and measuring BOD, the test substance exhibited < 1% after 5 days.

Unpublished, IUCLID(non-confidential), as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

9.2 BIODEGRADATION

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

METHOD

Method/guideline followed: Two laboratory anaerobic digesters, AD1 and AD2, were prepared by adding 180 ml treated well water (W13) (sampled from a well located under an area of coal-tar contaminated ground water and with a known alkylphenol content) and 2 ml of ferrous sulfide suspension to 250 ml serum bottles. Aliquots (20 ml) from a second well (P14) located 143 m down-gradient from the first were added to each digester as the inoculum. Both digesters were purged with oxygen-free nitrogen. Digester AD1 was operated in a semi-continuous addition mode for culture enrichment. At approximately 84 hour interval an aliquot of the solution was removed and replaced with fresh well water. AD2 was maintained for the eight weeks of the experiment with no additions or subtractions.

Methane concentrations in the headspace and alkylphenol concentrations in the aqueous phase were determined by gas chromatography.

Test type: anaerobic.

GLP (Y/N): no.

Year study performed: 1983.

Contact time: 8 weeks.

Inoculum: Well water (Well P14) from a zone of active microbiological activity.

Concentration: 0.37 mg/l.

RESULTS

Percentage degradation:

Digester AD1: 54% after 8 weeks

Digester AD2: 0% after 8 weeks.

Some of the other phenols present (e.g. 3-methylphenol) did undergo significant degradation in both digesters.

CONCLUSIONS

Analysis of a third well (P119) located 430 m down-gradient from Well W13 indicated that all the alkylphenols present in W13 underwent virtually complete anaerobic biodegradation after a sufficient time in the aquifer. The authors of the present study suggest that the design of the laboratory digesters may have inhibited complete biodegradation.

DATA QUALITY

Not a GLP study.

Adequate information is available regarding the experimental design and results. The latter confirm that alkylphenols have varying susceptibility to anaerobic biodegradation. The design of the laboratory digesters does not appear to model environmental processes adequately.

REFERENCES

Godsy, E.M. et al., Bull. Environ. Contam. Tox. 30, 261-268, (1983).

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

Bestimmung der Wirkung von Wasserinhaltsstoffen auf Fische, DIN 38412 Teil 15

Test type: Static

GLP: (N)

Year study performed: 1982

Species/strain: Freshwater fish, Leuciscus idus

RESULTS

NOEC 4.6 mg/l

LC0 (96hr) 10 mg/l

LC50 (96hr) 10-22 mg/l

LC100 (96hr) 21.5 mg/l

CONCLUSIONS

LC50 (96hr) value for 2,3,6-trimethyl phenol was found to be within the range of 10 to 22 mg/l.

DATA QUALITY

Information taken from IUCLID database.

Purity of the test substance: information not available in database.

GLP study: data not available.

REFERENCES

IUCLID Data Sheet

4. Ecotoxicity 4.1 Acute and Prolonged Toxicity to Fish

Study Ref.: BASF AG Ludwigshafen

OTHER – ADDITIONAL

In another flow-through fish acute toxicity study, using $Pimephales\ promelas$, a 96-hour LC50 of 8.2 mg/l was identified.

IUCLID (non-confidential), as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

10.2 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

IUCLID Identification Number: 0675

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 3.9 mg/l

ChV (30 day) 0.58 mg/l

ChV (90 day) 0.050 mg/l

Remarks: log Kow used 3.15 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 3.9 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

Scenedesmus-Zellvermehrungs-Hemmtest, DIN 38412 Teil 9, Bestimmung der Hemmwirkung von Wasserinhaltsstoffen auf Gruenalgen

Test type: Data not available

GLP: (N)

Year study performed: 1988

Species/strain: Scenedesmus subspicatus

RESULTS

EC20 (72hr) 15 mg/l

EC50 (72hr) 19 mg/l

EC90 (72hr) 24 mg/l

CONCLUSIONS

The EC50 (72hr) value for 2,3,6-trimethyl phenol was found to be 19 mg/l.

DATA QUALITY

Information was taken from IUCLID database. Purity of the test substance: information not available. GLP study: data not available.

REFERENCES

IUCLID Data Sheet

4. Ecotoxicity 4.3 Toxicity to Aquatic Plants

OTHER

Study Ref.: BASF AG Ludwigshafen

11.2 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 6.4 mg/l

ChV (96hr) 1.3 mg/l

Remarks: log Kow used 3.15 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 6.4 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

Directive 88/449/EEC, Method C.2

Test type: Data not available

GLP: (N)

Year study performed: 1988

Species/strain: Daphnia magna (Crustacea)

RESULTS

EC0 (24hr) 6.25 mg/l

EC50 (24hr) 12.6 mg/l

EC100 (24hr) 25 mg/l

CONCLUSIONS

The EC50 (24hr) value for 2,3,6-trimethylphenol was found to be 12.6 mg/l.

DATA QUALITY

Information taken from IUCLID database.

Purity of the test substance: information not available.

GLP study: data not available.

REFERENCES

IUCLID Data base

4. Ecotoxicity 4.2 Acute Toxicity to Aquatic Invertebrates

OTHER

Study Ref.: BASF AG Ludwigshafen

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

IUCLID Identification Number: 0675

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 2.5 mg/l

ChV (21day) 0.42 mg/l

Remarks: log Kow used 2.72 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be $2.5\ mg/l$.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

12.3 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

IUCLID Identification Number: 0675

The substances used in this test were purchased from different commercial sources and were not purified before testing (all > 95% purity).

METHOD

AFNOR (1974)

Acute Toxicity to Daphnia magna

Test type: Static

GLP: Data not available

Year study performed: 1987

Test substance preparation: The compounds were diluted with AFNOR (1974) reconstituted hard water for toxicity test (pH 7.8 –8.2; hardness, 200 mg/l expressed as CaCO₃). Phenol 2,3,6-trimethyl was dissolved directly in reconstituted water, which had been aerated for 1 day prior to use. Acetone was used as dispersent-solvent at a maximum concentration of 0.1 ml/l of reconstituted water (ISO,1980). Five daphnids were placed into the test tube and reconstituted water added to make the volume up to 10 ml.

Species: *Daphnia magna* Straus 1820 used in this experiment came from the IRCHA Laboratory and had been cultured parthenogenetically in the Pasteur Institute Laboratory.

Analytical procedures: None

Test details: Twenty-four-hour IC50 values for three trimethylphenols, phenol, o-cresol, m-cresol, p-cresol and six xylenols were determined for *Daphnia magna* under static conditions.

Statistical methods: Percentages of immobilisation (between 10 and 90% on the basis of total number of *Daphnia* per concentration) were recorded and plotted as a function of concentration on log-probit paper. The points obtained were fitted to a straight line from which the IC50 was read as the abscissa of the point corresponding to 50% immobilisation. Each chemical was assayed in duplicate with a minimum of three replicates.

RESULTS

Single series of geometrically spaced concentrations were used (0.1, 0.35, 1, 3.5, 10, 35, 100 and 350 mg/l) in both the preliminary and the definitive tests.

Observations were made after 24 hour exposure. Daphnids that were unable to swim within 15 minutes after stimulation by gentle agitation of the water were considered to be immobilised. The 24-hour IC50 value to *Daphnia magna* for 2,3,6 trimethylphenol was found to be 0.143 mmol/l with the 95% confidence interval (0.130 - 0.155).

CONCLUSIONS

The results of this test showed that addition of a methyl group to phenol increases the toxicity compared with that registered to phenol.

DATA QUALITY

Information taken from the test report.

GLP study: data not available.

REFERENCES

Acute Toxicity of Cresols, Xylenols and Trimethylphenols to Daphnia Magna Straus 1820

OTHER

Study Ref.: The Science of the Total Environment, 76(1988) 79-83

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

OECD Guideline No 401

GLP: (N)

Year study performed: 1986

Species/strain: Rat

Route of administration: Oral, gavage

RESULTS

LD50 >2000 mg/kg. Some mortality occurred at this dose level.

CONCLUSIONS

The LD50 of the test substance was found to be greater than 2000 mg/kg.

DATA QUALITY

Information taken from IUCLID database

Test performed to OECD guidelines but not GLP

REFERENCES

IUCLID Database

5.Toxicity, 5.1 Acute Toxicity, 5.1.1 Acute Oral Toxicity

OTHER

Study Ref: BASF AG Ludwigshafen

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6

OH

IUCLID Identification Number: 0675

METHOD

OECD Guideline No 471

The Salmonella Standard Plate Test (SPT) and Preincubation Test (PIT)

GLP: (N)

Year study performed: 1983

Species/strain: Salmonella typhimurium TA 1535, TA1537, TA 100 & TA 98

Concentrations tested:

Standard Plate Test

20-5000 µg/plate TA100, TA 98; 4-2500 µg/plate TA 1535, TA 1537

Preincubation Test: 4-5000 µg/plate all tester strains

With and without S-9 metabolic activation

RESULTS

Negative

CONCLUSIONS

The test material is not genetically active in the Ames *Salmonella typhimurium* assay In the presence or absence of S-9 metabolic activation.

DATA QUALITY

Information taken from IUCLID database

Test performed to OECD guidelines but not GLP

REFERENCES

IUCLID Database

5. Toxicity 5.5 Genetic Toxicity in Vitro

OTHER

Study Ref: BASF AG Ludwigshafen

15.2 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

2,3,6-Trimethylphenol

CAS No. 2416-94-6



IUCLID Identification Number: 0675

METHOD

OECD Guideline No 471

GLP: (Y)

Year study performed: 1987

Species/strain: Salmonella typhimurium, TA 1535, TA1537, TA100 & TA 98

Concentrations tested: $0.1500 \mu g/plate$ all tester strains

With and without S-9 metabolic activation

RESULTS

Negative

CONCLUSIONS

The test material is not genetically active in the Ames *Salmonella typhimurium* assay In the presence or absence of S-9 metabolic activation.

DATA QUALITY

Information taken from IUCLID database.

Test performed to OECD guidelines and to GLP.

REFERENCES

IUCLID Database

5. Toxicity 5.5 Genetic Toxicity in Vitro

OTHER

Study Ref: BASF AG Ludwigshafen

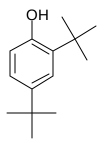
PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



METHOD

Method/guideline followed: ASTM D 1015.

GLP (Y/N): no.

Year study performed: 1994.

RESULTS

Melting point: 56.5°C.

CONCLUSIONS

The test substance has a melting point of 56.5°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Huels AG: Produktinformation "2, 4-Di-*tert*-butylphenol", Art. No. 001156, 01-Aug-1994 as cited in IUCLID database.

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1995.

RESULTS

Boiling point: 264°C at 101300 Pa (760 mm Hg).

CONCLUSIONS

The test substance has a boiling point of 264°C at 101300 Pa.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Huels AG: Sicherheitsdatenblatt "2, 4Di-tert-butylphenol", Version 04, 13.01.95 as cited in IUCLID database.

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1994.

RESULTS

Vapor pressure: ca. 1.0 Pa @ 20°C.

CONCLUSIONS

The test substance has a vapor pressure of ca. 1.0 Pa @ 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Huels AG: Produktinformation "2, 4-Di-tert-butylphenol", Art. No. 001156, 01-AUG-1994 as cited in IUCLID database.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 5.33.

CONCLUSIONS

The test substance has a log Kow of 5.33.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemic al Properties Prediction Program. KOWWIN v1.63.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1994.

RESULTS

Water solubility: 12 mg/l at 20°C.

CONCLUSIONS

The solubility of the test substance in water is 12 mg/l at 20°C.

DATA QUALITY

Not a GLP study.

Information taken from IUCLID database.

Purity of the test substance: information not available in database.

REFERENCES

Huels AG: Produktinformation "2, 4-Di-tert-butylphenol", Art. No. 001156, 01-AUG-1994 as cited in IUCLID database.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

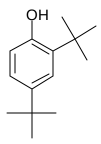
ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $49.1 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.61 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.61 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J. H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W. J., Reehl, W. F. and Rosenblatt, D. H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

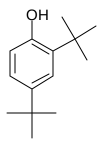
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 206
Data temperature (°C): 25
Water solubility (mg/l): 12
Vapor pressure (Pa): 1.0
Log Kow: 5.33
Melting point (°C): 56.5

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	1.75
Soil	95.6
Water	0.50
suspended sediment	0.066
fish	0.0054
Sediment	2.12

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days/weeks

Ultimate biodegradation in weeks/months.

This material is considered to be inherently biodegradable and would aerobically biodegrade slowly.

DATA QUALITY

Calculation method.

REFERENCES

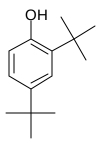
SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

9.2 BIODEGRADATION

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



METHOD

Method/guideline followed: ISO draft "BOD test for insoluble substances".

Test type: aerobic.

GLP (Y/N): no.

Year study performed: 1990.

Contact time: 28 days.

Inoculum: non-adapted activated sludge.

Concentration: 34.5 mg/l (related to test substance).

RESULTS

Percentage degradation: 2% after 28 days.

CONCLUSIONS

The test substance was not readily biodegradable under the test conditions.

DATA QUALITY

Not a GLP study.

Test concentration was higher than water solubility of test substance.

REFERENCES

Huels AG, report No. BO 90/5, 1990 (unpublished) as cited in IUCLID database.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

2, 4 Di- tert-butylphenol

CAS No. 96-76-4

METHOD

Static, DIN 38412 part 15

GLP: (unknown)

Year study performed: unknown

Species: Leuciscus idus (Golden orfe)

Exposure period: 48 hours

RESULTS

LC50 (48 hr) 1.8 mg/l

CONCLUSIONS

LC50 (48 hr) value for the test substance was found to be 1.8 mg/l.

DATA QUALITY

Not known, but IUCLID

REFERENCES

IUCLID (non-confidential), as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

10.2 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2, 4 Di- tert-butylphenol

CAS No. 96-76-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96hours, 30days, 90 days

RESULTS

LC50 (96hr) 0.27 mg/l

ChV(30day) 0.038 mg/1

ChV(90day) 0.008 mg/l

Remarks: log Kow used 5.33 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) value for the test substance was found to be 0.27 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

11.1 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2, 4 Di- tert-butylphenol

CAS No. 96-76-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.12 mg/l

ChV (96hr) 0.079 mg/l

Remarks: log Kow used 5.33 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) value for the test substance was found to be 0.12 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

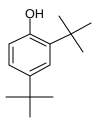
Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2, 4 Di- tert-butylphenol

CAS No. 96-76-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48hours, 21 days

RESULTS

LC50 (48hr) 0.48 mg/l

ChV (21day) 0.029 mg/l

Remarks: log Kow used 5.33 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) value for the test substance was found to be 0.48 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

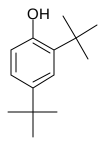
HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



No information is provided concerning the degree of purity of the material used in the test.

METHOD

Acute oral toxicity to rat

EPA CFR 163. 81-1, PSL -SOP No. 006

GLP: No data available

Year study performed: 1985

Species: Albino rats

Sex: Males and females

No of animals per sex per dose: 5 males and 5 females

Vehicle: 50% suspension in corn oil

Route of administration: Oral, gavage

Remarks:

A preliminary 7-day range finding test was conducted at five dose levels: 0.5, 1.0, 2.0, 4.0 and 8.0 g/kg bodyweight. Two rats were allotted to each dose level. Thereafter five groups, each consisting of 5 male and 5 female healthy albino rats were individually dosed by gavage at the following dose levels: 1.0, 1.5, 2.0, 2.5 and 3.0 g/kg/bodyweight.

After dosing the animals, their feed was returned and they were observed daily of mortality and other signs of gross toxicity for 14 days. Gross necropsies were performed on all mortalities and on all survivors at terminal sacrifice. At the end of the test period individual final body weights were recorded. The defined oral LD50 was calculated as by Litchfield – Wolcoxon Method of Probit Analysis (J. Pharmacology and Experimental Therapeutics 96: 99-115, 1949).

RESULTS

Acute oral LD50:

Total: 1.5 g/kg (95% confidence limits 1.2 - 1.9 g/kg)

Male: 2.1 g/kg (Estimated graphically)

Female: 1.3 g/kg (Estimated graphically)

30% mortality occurred in the test group dosed at 1.0 g/kg. Clinical signs included loss of appetite, hunched posture, inability to move and chromodacryorrhoea. Deaths occurred on days 4 and 5. The survivors recovered and returned to normal on Day 4. Prior to recovery, these rats were lethargic, and one had a bloody nasal discharge. All surviving animals gained weight and appeared healthy at terminal sacrifice

50% mortality (2 males, 3 females) occurred in the test group dosed at 1.5 g/kg within 6 days post dose. Preceding death symptoms included diarrhoea, loss of appetite, bloody ocular and nasal discharges and inability to move. Among the survivors, all had transient diarrhoea for several hours on the day of dosing. Other signs of toxicity prior to recovery included lethargy, loss of appetite in one case, accelerated breathing and hindered mobility. By Day 7 the surviving animals had regained all functions and gained weight. They continued to gain weight and remained active for the balance of the observation period.

40% mortality occurred in the test group dosed at 2.0 g/kg (2 males, 2 females). The two males and one of the females died on Day 4. The other female died on Day 6. Prior to death two of the animals were cyanotic and along with the rest showed variety of symptoms including loss of appetite, diarrhoea, chromodacryorrhoea, hunched posture, bloody nasal discharge, immobility and lethargy. By Day 7 all surviving animals appeared to recover after showing symptoms of lethargy, loss of appetite and no overt signs of intoxication. At terminal sacrifice, all survivors were active and all gained weight.

90% mortality (5 males and 4 females) occurred in the test group dosed at 2.5 g/kg. One rat died on Day 2; seven on Day 3; and one on Day 5. Preceding death animals showed toxic symptoms and signs that included bloody discharge from masal and ocular orifices, diarrhoea, inability to move and lethargy. Recovery was apparent by Day 8 and at terminal sacrifice the only surviving rat gained weight.

80% mortality was noted in the test animals dosed at 3.0 g/kg: one animal died on Day 2; 4 animals on Day 4; and 3 animals on Day 6. Signs and symptoms recorded before death included hematuria in two cases, cyanosis in four cases, diarrhoea in all cases, ataxia or unsteady gait in two cases, loss of appetite in all cases, bloody nasal and ocular discharges and lethargy. The surviving 20% also showed diarrhoea, loss of appetite, bloody discharges and lethargy. These two animals recovered and remained active following Day 9 of the test. One out of two animals had lost weight by Day 7 while the other remained at the same weight; both gained weight by terminal sacrifice.

Preliminary range finding

Sex	Dose level g/kg	Mortality day
M	0.5	-
F	0.5	-
M	1.0	-
F	1.0	-
M	2.0	4
F	2.0	-

M	4.0	3
F	4.0	3
M	8.0	3
F	8.0	3

Individual results

Dosage- 1.0 g/kg/bodyweight

Sex	Actual dose* ml/kg	Mortality day	Autopsy
M	0.40	Е	UR
M	0.40	Е	UR
M	0.40	Е	UR
M	0.40	Е	UR
M	0.40	Е	UR
F	0.39	4	BND; D; DL
F	0.39	5	DL; DGS
F	0.40	Е	UR
F	0.40	5	D; BND; DL
F	0.40	E	UR

Dosage- 1.5 g/kg/bodyweight

Sex	Actual dose* ml/kg	Mortality day	Autopsy
M	0.58	Е	UR
M	0.59	E	UR
M	0.59	6	LH; IH
M	0.59	E	UR
M	0.59	6	BND; BOD; D; IH; DK
F	0.58	Е	UR
F	0.58	3	BND; DL
F	0.59	6	D;BND; LH; IH
F	0.59	5	DGS; DL
F	0.59	Е	UR

Dosage- 2.0 g/kg/bodyweight

Sex	Actual dose* ml/kg	Mortality day	Autopsy
M	0.80	E	UR
M	0.80	4	BDN; D; PH; IH
M	0.81	Е	UR
M	0.83	Е	UR
M	0.83	4	BND; D; PH; DL;
			DGS; IH
F	0.75	E	UR
F	0.75	Е	UR
F	0.75	6	BND; PH; IH; DL
F	0.76	4	BND; D; LH; DGS;
			GIH
F	0.77	Е	DL

Dosage- 2.5 g/kg/bodyweight

Sex	Actual dose* ml/kg	Mortality day	Autopsy
M	0.92	Е	UR
M	0.94	3	PH; DL; DGS; IH
M	0.94	3	D; DGS; GIH; DL; SH
M	0.96	3	DGS; DL
M	0.97	5	DGS; DL; IH
F	0.95	3	BND; BOD; DL; DGS;
			IH
F	0.96	3	DGS; DL; SH
F	0.97	3	DGS; DL
F	0.97	2	D; IH; SH
F	0.97	3	DGS; DL; DK

Dosage- 3.0 g/kg/bodyweight

Sex	Actual dose* ml/kg	Mortality day	Autopsy
M	1.3	Е	UR
M	1.3	4	BND; D; DL; IH
M	1.3	6	BND; GIH; IH; DL
M	1.3	2	BND; D; GIH; DGS;
			LH; SH
M	1.3	4	BND; D; DGS; IH
F	1.2	Е	UR
F	1.2	4	BND; D; DH; DL
F	1.2	6	BND; GIH; DL; IH
F	1.2	4	PH; DL; DGS; IH
F	1.2	6	BND; DL; IH; GIH

^{*}The test material was uniformly mixed with corn oil and administered as a 50% w/w suspension by gavage.

Autopsy code

E Euthanized

UR Unremarkable

BND Bloody nasal discharge

D Diarrhea

PH Pulmonary hemorrhage

DL Discolored liver

IH Intestinal hemorrhage

DGS Distended gaseous stomach

GIH Gastro-intestinal hemorrhage BOD Bloody oral discharge

SH Splenic hemorrhage

DK Discolored kidney

LH Liver hemorrhage

DH Discoloration of the heart

CONCLUSIONS

LD50: 1.5 g/kg.

DATA QUALITY

Study conducted according to EPA guidelines. No data available on GLP.

REFERENCES

Report No.T-5051, Product Safety Labs, 725 Cranbury Road, East Brunswick, New Jersey 08816, USA.

OTHER - ADDITION

Another Rat Oral LD50 value of 2559-4128 mg/kg bw is in the IUCLID database for this compound. The methodology is cited as OECD 401.

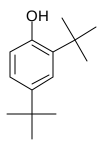
IUCLID (non-confidential, but unpublished), as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

13.2 ACUTE TOXICITY DERMAL IRRITATION

TEST SUBSTANCE

2,4-Di-tert butylphenol

CAS No. 96-76-4



METHOD

OECD Guidelines No. 404

GLP: (Y)

Year study performed: 1991

Species: New Zealand rabbits, White.

Sex: Male, females.

No of animals per sex per dose: One male, two females.

Vehicle: 0.5 g of the test substance.

Remarks: All the work was performed in compliance with the good laboratory practice; except that the concentration, homogeneity and stability of the test material preparations were not determined by analysis; and that this specific study may not have been subject to procedure inspection by the Quality Assurance Unit.

RESULTS

A single 4-hour, semi-occluded application of the test material to the intact skin of three rabbits produced well defined erythema and very slight to slight edema. Other skin reactions noted were hemorrhage of the dermal capillaries, blanching, light brown discoloration of the epidermis, crust formation and hardened light brown-colored scabs.

The test produced a primary irritation index 3.5 and was classified as a moderate irritant to rabbit skin according to the Draize classification scheme. No corrosive effects were noted.

CONCLUSIONS

The test material was also classified as irritant according to EEC labelling regulations. The symbol "Xi" and risk phrase R 38 "Irritating to skin" are therefore required.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

Project number: 47/1591, Safepharm Laboratories Limited, P.O Box 45, Derby, DE1 2BT,. U.K.

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Melting point: 36 - 37°C.

CONCLUSIONS

The test substance has a melting point range of 36 - 37°C.

DATA QUALITY

Not a GLP study.

Information taken from SIDS dossier.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Technical Bulletin FC:69:46:TB (Shell Chemicals)

Material Safety Datasheet, STIA, CH-4133 Pratteln, April 1989.

Both as cited in SIDS dossier for Phenol, 2,6 Bis(1,1-dimethyl).

2.1 BOILING POINT

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: not known.

RESULTS

Boiling point: 253°C at 10100 Pa (760 mm Hg).

CONCLUSIONS

The test substance has a boiling point of 253°C at 10100 Pa.

DATA QUALITY

Not a GLP study.

Information taken from SIDS dossier.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Technical Bulletin FC:69:46:TB (Shell Chemicals)

Material Safety Database, STIA, CH-4133 Pratteln, April 1989.

Both as cited in SIDS dossier for Phenol, 2,6-bis(1,1-dimethyl).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: gas saturation.

GLP (Y/N): yes.

Year study performed: 1990.

RESULTS

Vapor pressure: 0.0076 mm Hg at 20°C.

CONCLUSIONS

The test substance has a vapor pressure of ca. 0.0076 mm Hg at 20°C (1.01 Pa).

DATA QUALITY

GLP study.

Gas saturation is a recognised method described in EU, OECD and OPPTS guidelines.

REFERENCES

Vapor pressure of 2, 6-Di-t-Butylphenol. Unpublished report no. 56:89 from Ethyl Corporation, February 1990. As cited in SIDS dossier for Phenol, 2, 6-bis(1,1-dimethyl).

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: OECD 117, HPLC method.

GLP (Y/N): Yes.

Year study performed: 1992.

RESULTS

Log Kow: 4.5 at 24 - 26°C.

CONCLUSIONS

The test substance has a log Kow of 4.5.

DATA QUALITY

GLP study.

An acceptable test method has been used.

REFERENCES

Determination of the partition coefficient (n-octanol/water) of 2,6-bis(1,1-dimethylethyl)-phenol by HPLC. Unpublished RCC report no. 304424 from Sandoz Chemicals Ltd., March 1992 as cited in the SIDS dossier for Phenol, 2, 6-bis(1,1-dimethyl).

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: Generator Column Method in accordance with the Federal Register, Volume 50, Number 188, page 39265.

GLP (Y/N): yes.

Year study performed: 1988.

RESULTS

Water solubility: 4.11 mg/l at 25°C (pH 7), 3.99 mg/l at 25°C (pH 4), 4.69 mg/l at 25°C (pH 9).

CONCLUSIONS

The solubility of the test substance in water at pH 7 is 4.11 mg/l at 25°C and is unaffected by pH over the range pH 7 to 9.

DATA QUALITY

GLP study.

The generator column technique is an acceptable method.

REFERENCES

Springborn Life Sciences, Inc.: Determination of the water solubility of 2,6-Di-tert-butylphenol. Unpublished report number 88-07-2771, Study no. 10671-0887-6107-700, October 1988 as cited in the SIDS dossier for Phenol, 2, 6-bis(1,1-dimethyl).

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: 49.1×10^{-12} c m³/molecule-sec.

Half-life: 2.61 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.61 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

OTHER - ADDITIONAL

Photodegradation was reported and appears to be more important than biodegradation.

Freitag, D. et al, Ecotoxicology and Environmental Safety 6(1), 60-81 (1982).

SIDS of OECD High Production Volume Chemicals Programme (1994).

6.2 PHOTODEGRADATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: Determination of Indirect Photolysis, EPA TSCA Guideline 40 CFR 795-70.

Test type: indirect photolysis in simulated natural water.

GLP (Y/N): yes.

Year study performed: 1992.

Light source: natural light.

Light spectrum: that of natural light.

Light intensity: 67.5 to 476 footcandles.

Spectrum of substance (max lambda, max epsilon and epsilon 295): not stated.

Duration of test: the main test (phase III) lasted 29 hours (15 hours of daylight).

There were two main phases in the study. In the preliminary test (Phase II) approximate sunlight reaction rates were determined and an attempt made to differentiate between direct and indirect processes. From Phase III, the main study, net and direct photolysis rate constants were obtained.

Protocol deviations: all protocol deviations were minor and did not effect the results of the study except that HPLC-UV rather than HPLC-RAM was used for quantification of the test substance. This reduced the chance of detecting breakdown products.

RESULTS

Concentration of substance: 1.25 mg/l.

Temperature: *ca.* –0.2 to 26.5°C (samples were moved indoors each night to reduce the risk of freezing).

Indirect photolysis

Sensitizer (type): humic acid (in aqueous solution)

Concentration of sensitizer: 0.005% w/v.

Rate constants

Phase II

Indirect aqueous photolysis: 2.94 day⁻¹.

Direct aqueous photolysis: 2.16 day⁻¹.

Phase III

Net photolysis: 6.90 day⁻¹.

Direct aqueous photolysis: 15.17 day⁻¹.

Percentage degradation: photolytic half-life under environmental conditions 2.41 hours.

Breakdown products: no volatile breakdown products were observed.

CONCLUSIONS

2,6-Di-tertiary butylphenol is marginally susceptible to indirect photolysis in natural waters. The net photolytic half-life under environmental conditions was found to be 2.41 hours

DATA QUALITY

GLP study.

There was significant loss of the test substance from controls. This is believed to have been due to sorption onto glass.

REFERENCES

Springborn Laboratories Inc. 790 Main Street, Wareham, Massachusetts 02571, USA. Test report no. 92-1-4065, 5 August 1992.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W. J., Reehl, W. F. and Rosenblatt, D. H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

OTHER - ADDITIONAL

Considered unstable in water.

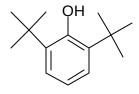
Unpublished study in IRPTC Data profile as cited in "An Initial Review of Potential Environmental Risks from Alkylphenols other than Nonylphenol," produced for the UK Environment Agency by WRc-NSF.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:206Data temperature (°C):25Water solubility (mg/l):4.11Vapor pressure (Pa):1.01Log Kow:4.5Melting point (°C):36.5

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	25.6
Soil	70.3
Water	2.51
suspended sediment	0.049
Fish	0.0040
Sediment	1.56

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: EEC, L 251 Vol. 27 (19.09.84) 84/449/EEC.C.5.

Test type: Ready biodegradability (aerobic) by the Modified Stürm Test.

GLP (Y/N): no.

Year study performed: 1990.

Contact time: 28 days.

Inoculum: Bacteria collected from activated sludge.

Concentrations: 10.5 and 20.2 mg/l of the test substance (these concentrations were in excess of the water solubility of the test substance; nonylphenol was therefore used as an emulsifier).

RESULTS

Percentage degradation

Lower concentration: 4% after 28 days.

Higher concentration: 1% after 28 days.

Results calculated based on measured CO₂ formation vs. theoretical carbon content of the test solution.

CONCLUSIONS

2,6-Di-tertiary butylphenol was not readily biodegradable under the conditions of this test.

DATA QUALITY

Not a GLP study.

REFERENCES

CIBA-GEIGY Ltd., Basle, Switzerland. Test report no. 894579, study completed 19.02.90. Title: Report on the test for ready biodegradability in the Modified Stürm Test of IRGANOX L 108.

9.2 BIODEGRADATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Method/guideline followed: TSCA Test Guideline 796.3140.

Test type: Aquatic biodegradation under anaerobic conditions.

GLP (Y/N): yes.

Year study performed: 1991.

Contact time: 56 days.

Inoculum: Primary sludge from a wastewater treatment plant.

Concentration: 63 mg/l.

RESULTS

Percentage degradation: none.

The test concentration was considerably in excess of the water solubility of 2,6 di-tertiary butylphenol. There will therefore have been undissolved and sorbed material present in the test system. Some material may also have been present in the headspace of the test system.

CONCLUSIONS

2,6-Di-tertiary butylphenol was not biodegradable under the conditions of the study.

DATA QUALITY

GLP study.

REFERENCES

Springborn Laboratories Inc., 790 Main Street, Wareham, Massachusetts 02571, USA. Test Report no. 92-2-4105 (unpublished), (8/5/92).

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

Purity of the test substance: information not available.

METHOD

Acute Toxicity to Fish

Directive 84/449/EEC, Method C1

Test type: Static

GLP: Data not available

Year study performed: 1990

Species: Zebra-fish (Brachydanio rerio)

Analytical procedures: Data not available.

Remarks: Small parts of the test substance were floating at the surface of the tanks at 0 and 96-hour exposure.

Exposure period: 96 hours

Test conditions:

- dechlorinated tap water (carbon filter),
- water hardness 196 mg CaCO₃/l
- temperature 23°C, pH 8.0 8.1, O₂ 100 94% (0-96hr measurements)

Statistical methods: The LC50 values were calculated according to Berkson, Jasa 48 (1953), 569-599.

LC-values were graphically determined on gausso-logarithmic probability paper.

RESULTS

Nominal concentrations: 1.0, 1.8, 3.2, 5.8, 10 and 18 mg test substance /l

All results are expressed in mg/l and are base on nominal concentrations.

Calculated LC50 values with 95% confidence limits:

LC50 (24hr), 9.7 mg/l (7.8-12)

LC50 (48hr), 7.6 mg/l (6.5-8.9)

LC50 (72hr), 7.6 mg/l (6.5-8.9)

LC50 (96hr), 7.6 mg/l (6.5-8.9)

Graphically determined LC50 value:

LC50 (96hr), 7.6 mg/l

Observed LC50 values:

LC0 (96hr) in test 5.8 mg/l

LC100 (96hr) in test, 10 mg/l

Controls: Mortalities in blank: 0%

CONCLUSIONS

Calculated LC50 (96hr) value for 2,6-Di-tert-butylphenol was found to be 7.6 mg/l.

DATA QUALITY

Information taken from the test report.

Purity of the test substance: information not available.

GLP study: no data available.

REFERENCES

Test Report on the Acute Toxicity Test of IRGANOX L 108 to Zebra-Fish (*Brachydanio rerio*). Ciba-Geigy Ltd, AD-Division, CH-4002, Basel, Switzerland.

OTHER

Study Ref.: 894

10.2 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

The sample used in this study was identified to be 99.88% active ingredient (A.I.).

METHOD

Protocol for Conducting a Flow-through Acute Toxicity Test with 2,6-Di-tert-butyl phenol and Freshwater Fish (*Oncorhynchus mykiss*) RBTFA/DTBP.

Test type: 14 day Flow-through exposure

GLP: (Y)

Year study performed: 1989

Species: Rainbow trout (Oncorhynchus mykiss), obtained from a commercial supplier in California.

4.3 ppm.

Analytical monitoring: All replicate treatment levels and the controls were analysed for ¹⁴C-2,6-DTBP concentrations prior to the start of the definitive test. During the in-life phase of the definitive study, water samples were removed from both replicate test solutions of each treatment level and the controls on test days 0, 4, 6, 8, 12 and 14 for analysis of ¹⁴C-2,6-DTBP. Each exposure solution sample was collected from the approximate midpoint of the aquarium with a volumetric pipette. Immediately after sampling, all samples were centrifuged for 25 minutes at 2000 rpm. This procedure was sufficient to remove suspended particles (≥ 0.45 microns) and absorbed test substance. In addition, quality assurance QA blind samples were prepared at each sampling interval and remained with the set of exposure solution samples through the analytical process. The QA samples were prepared in dilution water at 2,6-DTBP concentrations unknown to the analyst. Results of this analysis were used to judge the precision and quality control maintained during the analysis of exposure solution samples. All samples were analysed for ¹⁴C-2,6-DTBP using radiometric procedure.

Exposure period: 14 days

Statistical methods:

The mean measured concentrations tested (day 0, 6, 8, 12 and 14) and the corresponding mortality data derived from the toxicity test were used to estimate the median lethal concentrations (LC50) and 95% confidence interval at each 24-hour interval of the exposure period. If at least one test concentration caused mortality of greater or equal to 50% of the test population, then a computer program (Stephan, 1977, 1982) was used to calculate the LC50 values and 95% confidence intervals. Three statistical methods were available in the computer program: moving average angle analysis, probit analysis and nonlinear interpolation with 95% confidence intervals calculated by binomial probability. The selection of reported LC50 values and 95% confidence intervals was based upon an examination of the data base and the results of the computer analysis. Selection criteria included the establishment of a concentration-effect relationship (mortality), the number of concentrations causing partial responses and the span of responses bracketing the LC50 value. If two or more statistical methods produced acceptable results, then the method which yielded the smallest 95% confidence intervals was selected. The No Observation Effect Concentration (NOEC) was also determined.

Protocol deviations:

The protocol states that the exposure solution samples will be passed through a 0.45 micron filter prior to analyses. During the study the exposure solution samples were centrifuged for 15 minutes at 2000 rpm. This procedure was sufficient to remove particles (0.45 microns) and sorbed substances. Preliminary investigations performed at SLS established that filtering the exposure solutions removed soluble test material.

The protocol states that the exposure solution temperatures will be maintained at $12 \pm 1^{\circ}$ C. Results of the daily measurements in each vessel demonstrated that the solution temperatures ranged from 11 - 12° C. Continuous measurements in one replicates vessel resulted in a temperature range of $10 - 12^{\circ}$ C.

The protocol states that the concentration of solvent in the highest test concentration will not exceed $100~\mu l$. During the study the concentration of solvent in the highest treatment level was $446~\mu l$ acetone/l. Following consultation with EPA and the Study Sponsor, it was decided to exceed the recommended solvent limitation to allow the preparation and maintenance of consistent exposure concentration of 2.6~DTBP.

The protocol states that the TOC of the dilution water shall be measured at the beginning of the toxicity test and should not exceed 2.0 mg/l. Due to the analytical instrumentation, the samples removed at the test initiation could not be analysed. Historical values for TOC content of the dilution water averaged 4.3 ppm.

The protocol states that each replicate exposure solution will be sampled and analysed on test days 0, 4, 8, 12 and 14. Due to the lower than expected recoveries for the QA samples on day 4, additional samples were analysed on day 6. After review of the analytical data and the records describing the function of the diluter system, it was established that the results of the analysis of the solution on day 4 were not representative of the concentration of 2,6-DTBP in the exposure vessels. Therefore, mean measured concentrations for this study were based on the analyses performed on day 0, 6, 8, 12 and 14.

The protocol states that the dissolved oxygen concentration must exceed 90% at the initiation of the study and is not allowed to fall bellow 8.2 mg/l for the duration of the test. At test initiation, the dissolved oxygen concentration was 89% of saturation in one replicate (B) of the 1.5 mg A.I./l (nominal) treatment level. The dissolved oxygen concentration was equal to 90% of saturation in two additional replicate aquaria (replicate A of the 1.5 mg A.I./l nominal test concentration and replicate B of the 0.98 mg A.I./l nominal test concentration). In all remaining test aquarium, the dissolved oxygen concentration exceeded 90% of saturation at test initiation. In addition, on test days 13 and 14, the

dissolved oxygen concentration fell below 8.2 mg/l. However, the dissolved oxygen concentration never fell below 7.5 mg A.I./l (69% saturation) throughout the exposure period.

RESULTS

Nominal concentrations: 1.5, 0.98, 0.63, 0.41 and 0.27 mg A.I./l

Mean measured concentrations: 1.0, 0.66, 0.43, 0.28 and 0.21 mg A.I./l

All results are expressed in mg A.I./l

LC50 values for exposure days 1-6 were empirically estimated as being greater than the highest mean measured concentration: LC50 > $1.0 \,\mathrm{mg}$ A.I./l

LC50 values with 95% confidence intervals, calculated by moving average angle analysis:

LC50 (7-day), 0.89 mg A.I./l (0.78 – 1.1)

LC50 (8-day), 0.87 mg A.I./l (0.77 – 1.0)

LC50 (9-day), 0.83 mg A.I./l (0.75 - 0.93)

LC50 (10-day), 0.83 mg A.I./I (0.75 – 0.93)

LC50 (11-day), 0.81 mg A.I./l (0.70 – 1.0)

LC50 (12-day), 0.74 mg A.I./l (0.66 - 0.95)

LC50 (13-day), 0.89 mg A.I./l (0.66 - 0.85)

LC50 (14-day), 0.74 mg A.I./l (0.66 - 0.85)

Following 14 day exposure, 95% mortality was observed at the highest test concentration. Mortality of 20, 10, 65 & 20% was observed at the remaining lower treatment levels.

Due to the inversion of the concentration–effect relationship (65% mortality at 0.28 mg A.I./l and 10% mortality at 0.43 mg A.I./l), the reported 95% confidence intervals may not accurately reflect this statistic.

NOEC (14 day) < 0.21 mg A.I./l

Remarks:

Biological observations were made twice daily during the definitive study. Mortalities were recorded and removed from each aquarium every 24 hours during the exposure.

Throughout the exposure period, a small amount of precipitate was observed in the diluter system's mixing chamber. However, no undissolved 2,6-DTBP (e.g., precipitate film on solution surface) was observed in any of the exposure vessels during the study. The diluter system functioned properly.

Mean measured concentrations (mg A.I./l) and cumulative mortalities (%) during the 14 day flow-through study:

0.21 mg A.I./l; 0% (day 1), 5% (day 2 & 3), 10% (day 4), 20% (days 5 to 14)

0.28 mg A.I./l; 5% (day 1), 25% (day 2), 30% (day 3), 65% (days 4 to 14)

0.43 mg A.I./1; 5% (day 1), 10% (days 2 to 14)

0.66 mg A.I./1; 0% (day 1), 10% (day 2), 15% (day 3 to 10), 20% (days 11 to 14)

1.0 mg A.I./1; 0% (day 1), 5% (day 2), 20% (day 3), 35% (day 4), 40% (day 5), 45% (day 6), 65% (day 7), 70% (day 8), 80% (days 9 to 11), 95% (day 12 to 14)

Solvent control; 0% (days 1 to 14)

Control; 0% (days 1 to 14)

CONCLUSIONS

Based on a comparison of the 7- and 14- day LC50 values (0.89 and 0.74 mg A.I./l respectively), 2,6-DTBP does not appear to be chronically toxic to rainbow trout (*Oncorhynchus mykiss*).

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Acute Toxicity of 2,6-Di-*tert*-butyl phenol (2,6-DTBP) to rainbow trout (*Oncorhynchus mykiss*), during a 14 day study under flow-trough conditions. Springborn Laboratories Inc, 790 Main Street, Wareham, Massachusetts 02571, USA.

10.3 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.90 mg/l

ChV (30 day) 0.13 mg/l

ChV (90 day) 0.019 mg/l

Remark: log Kow used 4.48 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.90 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission.

10.4 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

Purity of the test substance given as 99.88%.

METHOD

Acute Toxicity to Fish

Test type: Flow-through

GLP: (Y)

Year study performed: 1989

Species: Rainbow trout (Oncorhynchus mykiss)

Solvent: Acetone

Exposure period: 4 to 14 days

Concentration:

0.74 - 1.0 mg/l

Doses of 0.89 mg/l were also tested for 7 days.

RESULTS

LC50 (96hr) > 0.1 mg/l

LC50 (7days) 0.89 mg/l

LC50 (14days) 0.74 mg/l

CONCLUSIONS

TheLC50 (96hr) for the test substance was found to be greater than 0.1 mg/l.

DATA QUALITY

GLP study

Information taken from the OECD/SIDS data set.

REFERENCES

Primary reference:

Ethyl Corporation. Unpublished Report-Ethyl Corporation, 89-05-2948, (1989)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref.: IRPTC Data Profile

10.5 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

Purity of the test substance: unknown.

METHOD

Acute Toxicity to Fathead Minnow

Test type: Flow-through

GLP: (Y)

Year study performed: 1989

Species: Fathead minnow (Pimephales promelas)

Solvent: Acetone

Exposure period: 4 to 14 days

Concentration: 1.0 - 1.4 mg/l

Doses of 1.1 mg/l were also tested for 7 days.

RESULTS

LC50 (96hr) 1.4 mg/l

LC50 (7days) 1.1 mg/l

LC50 (14days) 1.0 mg/l

CONCLUSIONS

TheLC50 (96hr) for the test substance was found to be 1.4 mg/l.

DATA QUALITY

GLP study

Information taken from the OECD/SIDS data set.

REFERENCES

Primary reference:

Ethyl Corporation. Unpublished Report-Ethyl Corporation, 88-12-2867, (1989)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref.: IRPTC Data Profile

10.6 ACUTE TOXICITY TO FISH - ADDITION

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

Purity of the test substance: unknown.

METHOD

Acute Toxicity to Zebrafish – OECD Guideline 203

Test type: Static

GLP: (N)

Year study performed: 1987

Species: Zebrafish (Brachydanio rerio)

Solvent: Acetone

Exposure period: 24 to 96 hours

Concentration: 10-24 mg/l

Doses of 13 and 15 mg/l were also tested for 48 and 72 hours.

RESULTS

LC50 (24 hr) 24 mg/l

LC50 (48 hr) 15 mg/l

LC50 (72 hr) 13 mg/l

LC50 (96 hr) 10 mg/l

Values are based on nominal concentrations; small oil droplets were observed on the surface of the test solutions.

CONCLUSIONS

TheLC50 (96hr) for the test substance was found to be 10 mg/l.

DATA QUALITY

Not a GLP study

Information taken from the OECD/SIDS data set.

REFERENCES

Primary reference:

Ciba-Geigy. Unpublished Report-Ciba Geigy, 87-40-58, (1987)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref.: IRPTC Data Profile

11.1 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

The test substance used in this report was 99.88% active ingredient (A. I.), a white gelatinous, viscous liquid received from Ethyl Corporation. After storage at room temperature, the liquid became a crystalline solid. The solid form of 2,6-Di-*tertiary* butylphenol was used in this study.

The radiolabelled material, a yellow-green-colored crystalline solid, was received as [Ring-U-¹⁴C] 2,6-Di-tertiary butylphenol (0.89 mCi/mmole) in three vials. Prior to testing, the 2,6-Di-tertiary butylphenol was stored under refrigerated conditions at 4°C.

METHOD

"96-Hour Toxicity Test with Freshwater Alga, *Selenastrum capricornutum*". EPA TSCA Guideline 797.1050 as amended on 20th May 1987 (U.S. EPA, 1987).

Test type: Static

GLP: (Y)

Year study performed: 1988

Species/strain:

Freshwater green alga *Selenastrum capricornutum*, obtained from Carolina Biological Supply Company, Burlington, North Carolina and maintained in stock culture at SLS.

Element basis:

Reduction in cell density (relative to the control).

Test duration:

96 hours (17 –21 October 1988)

Analytical monitoring:

At 24, 48, 72 and 96 hours, cells counts were conducted for each replicate vessel using a hemacytometer and microscope. When possible, approximately 400 cells per replicate were counted to provide \pm 10% accuracy at the 95% confidence level. At test initiation samples were removed from each of the 500 ml volumetric flasks of each treatment level and controls for analysis. At test

termination 0.5 ml was removed from each replicate of the lowest test concentration which completely inhibited algal growth, or of the highest test concentration in which growth inhibition occurred. The 1.5 ml of combined sub-sample was diluted with sterile MBL medium (pH adjusted to 7.5) to prepare a subculture with a 2,6 DTBP concentration equal to the highest test concentration in which growth inhibition was not observed. This subculture was used to determine if the effects of the test material on algae were algistatic (in which case cells would resume growth in the subculture) or algicidal (no growth would occur in the subculture). The subculture was incubated under the test conditions for up to 9 days or until growth was observed (as determined by daily cell counts). At the test termination, after the 0.5 ml sub-samples had been taken for sub-culturing, the contents of the three replicates of each test concentration were combined for 2,6-DTBP analysis. The pooled test solutions were filtered through 0.45 µm membrane filters, and both filters and filtrates were radio-assayed for 2,6-DTBP.

Statistical methods:

EC10, EC50 and EC90 values and their 95% confidence limits were determined by linear regression of response (percent reduction of cell density, as compared to controls) vs. nominal exposure concentration over the range of test concentrations where a clear exposure-response relationship was observed. Four linear regressions were estimated based on untransformed data, untransformed response vs. logarithm-transformed concentration, probit-transformed response vs. untransformed concentration and probit-transformed response vs. logarithm-transformed concentration. The regression that best fitted the data was selected based on the highest coefficient of determination (r). This regression equation was then applied to estimate the EC10, EC50 and EC90 values and their 95% confidence limits, using the method of inverse prediction (Sokal and Rohif, 1981). A computer program developed and validated at SLS was used to assist in these computations.

Deviations to protocol:

According to protocol the algae cultures are transferred into fresh medium regularly to provide six- to eight-day-old inoculum cultures. For this test the algae, which had been cultured in MBL medium were transferred into fresh medium five days prior to testing.

According to the protocol, the algal cultures and test cultures are maintained at a temperature of $24 \pm 1^{\circ}$ C. In this test, temperature ranged from 22 to 24° C.

The protocol states that at test termination, a 0.5 ml sample of culture is taken from each replicate of the lowest test concentration that completely inhibited algae growth to differentiate between algistatic and algicidal effects. During this study 1.0 ml from each replicate of an additional concentration, which severely inhibited growth, was taken as well.

Test conditions:

An inoculum of *Selenastrum capricornutum* cells calculated to provide 1 x 10^4 cells/ml (95% confidence interval 0.74 -1.26 x 10^4 cells/ml) was aseptically introduced to each flask. All flasks were than placed in an environmental chamber where the temperature was maintained at $22-24^{\circ}$ C. The flasks were impartially placed on the shaker, at the shaking rate of 100 rpm and were returned to the same position on the shaker after each daily observation. Lighting was provided continuously at intensity of 5,000 lux at the solution surface.

The MBL test medium used to prepare the exposure solutions was formulated in the same manner as the culture medium (excluding Na₂EDTA). The pH of this medium was adjusted to 7.5 ± 1 with 0.1 N hydrochloric acid.

Controls used in this test were controls (MBL medium) plus solvent (acetone) controls. The flask contained 50 µl of acetone (the greatest amount of solvent). Replicate sterile 125-ml flasks (3 per treatment level and the controls) were conditioned by rising with the appropriate test solution. 50 ml of

the appropriate test solution were then placed in each replicate flask. Control flasks were prepared and maintained under the same conditions as test flasks containing 2,6 DTBP. The opening of each flask was covered with a stainless steel cap to permit gas change.

pH at test initiation was 7.4 and at test termination pH ranged from 7.2 to 9.8.

RESULTS

Based on the results of a range-finding toxicity test, the nominal concentrations were 0.64, 1.3, 2.5, 5.0 and 10 mg/l.

Measured concentrations at test initiation ranged from 34 to 48% of nominal concentrations. Concentrations at 96 hours were extremely low, near the limits of detection of the analysis.

Based on mean of 0-hour and 96-hour analysis, the concentrations of 2,6-DTBP in this test were 0.11, 0.24, 0.51, 1.23 and 2.17 mg/l. In general, cell counts were relatively consistent among replicates at each test concentration. The exposure-response relationship was clearly defined, with the threshold for inhibition between mean measured concentrations of 1.23 and 5.1 mg/l. For counts made at 24, 72 and 96 hours, there were no significant differences in cell densities between controls and solvent controls, and the two sets of controls were combined for EC50 calculations. For counts made at 48 hours, cell densities were significantly higher in the solvent controls than in the controls, and EC50 values were based on the solvent controls.

Results based on mean measured concentrations with 95% confidence limits:

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24-hour EC10 value, 0.36 mg/l (0.16-0.80 mg/l)
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24-hour EC50 value, 0.86 mg/l (0.39-1.9 mg/l)

24-hour EC90 value, 2.2 mg/l (0.98-5.3 mg/l)

48-hour EC10 value, 0.16 mg/l (0.06-0.37 mg/l)

48-hour EC50 value, 0.50 mg/l (0.20-1.2 mg/l)

48-hour EC90 value, 1.6 mg/l (0.66-4.1 mg/l)

72-hour EC10 value, 0.29 mg/l (0.15-0.54 mg/l)

72-hour EC50 value, 0.51 mg/l (0.28-0.96 mg/l)

72-hour EC90 value, 0.96 mg/l (0.52-1.8 mg/l)

96-hour EC10 value, 0.18 mg/l (0.05-0.57 mg/l)

96 hours EC50 value, 0.56 mg/l (0.17-1.9 mg/l)

96-hour EC90 value, 1.7 mg/l (0.56-6.5 mg/l)

The control response was satisfactory.

The amount of filterable 2,6-DTBP at 96 hours could not be calculated precisely because the volume of culture filtered was not measured, but the total amounts of 2,6-DTBP on the filters were very low, ranging from 0.56 μ g to 6.12 μ g. Assuming that the volume filtered was between 50 and 100 ml, the amounts of 2,6-DTBP recovered on the filters correspond to a very small fraction of 2,6-DTBP initially present – i. e. filterable 2,6-DTBP was 0.0056 – 0.011 mg/l in the nominal 0.64 mg/l solution, and 0.061 – 0.12 mg/l in the nominal 10 mg/l solution. The loss of the test substance from the test solutions between 0 and 96 hours is unexplained. It is possible that the unaccounted 2,6-DTBP was present as undissolved material that settled to the bottom of the test vessels and was not included in the samples removed for filtration. Very little of the test material remained in solution at the termination of the study. The TSCA guidelines (797.1050) acknowledge that "For Chemicals with low solubility under test conditions, it may not be possible to quantitatively determine the EC50 value" (U.S. EPA, 1985).

CONCLUSIONS

Since it was established during this study that 2,6-DTBP was unstable in test solutions, it was not possible to quantitatively determine an EC50 value. The test substance caused reduction in cell density, relative to control in the toxicity test with the freshwater green alga *Selenastrum capricornutum*.

The re-growth of the algal subcultures indicated that the effects of 2,6-DTBP on *Selenastrum capricornutum* were algistatic, not algicidal.

DATA QUALITY

GLP study

Information taken from the test report.

REFERENCES

Toxicity Test with Freshwater Green Alga *Selenastrum capricornutum*, Springborn Laboratories, Inc., Environmental Sciences Division, 790 Main Street, Wareham, Massachusetts 02571, USA.

OTHER

Study Ref.: 88-11-2846

11.2 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

The test material was received in radiolabelled and nonradiolabelled form. A sample of 2,6-DTBP, a pale yellow crystalline solid identified by the sponsor to be 99.86% active ingredient, was received from Ethyl Corporation, Lousiana. This material was stored at room temperature $\sim 20^{\circ}\text{C}$ in the dark until use in the toxicity test.

METHOD

"96-Hour Acute Toxicity Test with Freshwater Alga, *Selenastrum capricornutum*". EPA TSCA 797-1050", as amended on 20th May 1987 (U.S. EPA, 1987).

Test type: Static

GLP: (Y)

Year study performed: 1991

Species/strain:

Freshwater green alga *Selenastrum capricornutum*; obtained from Carolina Biological Supply Company, North Carolina and maintained in stock culture at SLI.

Element basis:

Reduction in cell density after 24, 48, 72 and 96 hours of exposure (relative to the control).

Test duration:

96 hours (17-21 June 1991)

Analytical monitoring:

At test initiation, samples were collected form the volumetric flasks used to prepare the test solutions and analysed for 2,6-DTBP. At each 24-hour interval, three replicates from each test concentration, controls and solvent control were selected impartially. Aliquots of approximately 50 ml were removed from each of the three replicates and analysed. One of the nine 1.0 mg/A.I./l flasks (without algal inoculum) was sampled at test initiation for 2,6-DTBP analysis and the results used to determine the loss of 2,6-DTBP from solution in the absence of algae. All solutions were analysed using HPLC-UV (high performance liquid chromatography – ultraviolet detection HPLC-RAM (high performance liquid chromatography – radiometric) detection and LSC (liquid scintillation counting). A timeweighted average concentration was calculated for each set of test solutions at each sampling interval.

Each time-weighted average was calculated by averaging measurement for that interval with measurements made at the previous intervals.

Statistical methods:

The first stage of statistical analysis was comparison of control cell densities with solvent control densities for each sampling interval, using Student's T-test (Sokal and Rohlf, 1981). EC50 values based on initial measured concentrations and on time-weighted average concentrations were calculated from cell densities observed after 24, 48, 72 and 96 hours of exposure. For each observation period, the EC10, EC50 and EC90 values and their 95% confidence limits were determined by linear regression of response vs. the initial time-weighted average concentration over the range of test concentrations where a clear exposure-response relationship was observed. Four linear regressions were estimated based on a) untransformed data, b) untransformed response vs. logarithm-transformed concentration, c) probit-transformed response vs. untransformed concentration and d) probittransformed response vs. logarithm-transformed concentration. The regression that best fitted the data was selected based on the highest coefficient of determination (r^2) . This regression equation was then applied to estimate the EC values and their 95% confidence limits, using the method of inverse prediction (Sokal and Rohlf, 1981). The No Observed Effect Concentration (NOEC) based on initial measured concentrations and time-weighted average concentrations was determined using one-way analysis of variance (Sokal and Rohlf, 1981) and Dunnett's Test (Dunnett 1955, 1964) if all groups had equal numbers of replicates (i.e., if control data were not pooled), or using Bonferroni's Test (Weber, et al., 1989) if the groups had unequal numbers of replicates (i.e., if control data were pooled). Before conducting the analysis of variance, data were checked for normality using the Chi-Square test (Sokal and Rohlf, 1981) and for homogeneity of variance using the Hartley test (Neter et al., 1985). If the assumptions of normality and homogeneity of variance were not met, the NOEC was determined using the non-parametric Kruskal-Wallis Test (Sokal and Rohlf, 1981).

Deviations to protocol:

The protocol states that cell counts will be determined daily in the reference control solution. The reference control solutions were inadvertently not counted on day 1.

The protocol states that the analytical samples will be centrifuged prior to analysis. During the study, centrifugation of the samples was not needed because algal densities were low and did not interfere with direct injection HPLC analysis.

The protocol states that the method of choice for measurement of 2,6-DTBP is HPLC using direct aqueous injection followed by RAM. During the study HPLC using direct aqueous injection and UV detection was performed at each interval in addition to HPLC-RAM used for analysis on days 0 and 1 and which indicated that no radiolabelled degradates were detectable in the test solutions. LSC method was used to detect total ¹⁴C-residues on days 2,3 and 4. The test concentrations fell below the detection limit of HPLC-RAM.

The protocol states that the mass balance will be reported. Mass balance of test material was not determined during this study.

The protocol states that control solutions and QC spikes will be analysed at each interval. QC samples were not analysed by LSC on days 2 through 4. However LSC results are similar to HPLC-UV results.

The protocol states that no EDTA is added to the test medium. Based on a request from U.S. EPA, Na₂EDTA. H₂O was added to the test medium at a concentration of 0.38 mg/l. The protocol states that each test concentration will be analytically confirmed.

These deviations did not affect the results of the test.

The culture medium used was Marine Biological Laboratory (MBL) medium prepared with distilled water. The pH of this medium was adjusted to 7.5 ± 1 . Stock cultures were grown in 125 ml glass flasks containing 50 ml of medium.

Test conditions:

During the 96 hour test, the temperature was maintained at $23 - 25^{\circ}$ C. The flasks shaking rate was maintained at 100 rpm. Lighting was provided continuously by a combination of fluorescent light. The light was 300 to 400 f. candles.

The stock cultures were main tained under test conditions (shaking rate of 100rpm, temperature of 23 - 25°C, continuous illumination at the surface of approximately 375 to 500 f. candles).

The control solution was prepared containing algal growth medium only. In addition, a solvent control solution was prepared containing 0.10 ml/l acetone, equal to the concentration of acetone present in each test solution.

pH at test initiation was 7.5 and at termination pH ranged from 7.9 to 9.7.

A radiolabelled stock solution was prepared by quantitative transfer of the entire amount of ¹⁴C-DTBP (1.159 g) in one vial into a 25-l volumetric flask and diluting to volume with acetone. Radiochemical purity of this stock solution was determined to be 97.4% using HPLC with RAM detection. A nonradiolabelled stock solution of 158 mg A.I./ml was prepared by weighing 1.5824 g (1.5802 g A.I.) 2,6-DTBP into a sterile 10-ml flask and diluting to volume with acetone. A second nonlabelled stock solution of 58.1 mg A.I./ml was prepared. Seventy-two test vessels (twelve replicates for each test concentration, control and solvent control) were conditioned by rinsing with the appropriate test solution, control solution or solvent control solution. Nine additional flasks, identical to the exposure vessels, were prepared with 100 ml of the 1.0 mg A.I./l (nominal) solution for determination of 2,6-DTBP concentrations in the absence of algae.

Guideline requirements and modifications:

Requirement: Test solution volume does not exceed 50% of the flask volume. Modification: Test solution volume was 100 ml maintained in 125-ml Erlenmeyer flask.

Requirement: Vessel caps which allow gas exchange. Modification: Special exposure systems were use to minimise loss of 2,6-DTBP from the test solutions. Exposure vessels were 125-m1 glass Erlenmeyer flasks with teflon-coated stoppers.

Requirement: Provide an additional exposure at termination of the 96-hour exposure to determine algicidal/algistatic effects of the test material. Modification: Algicidal/algistatic properties of the test material were not investigated at termination of the 96-hour exposure. The exposure concentrations had declined to levels where this investigation was not meaningful.

Requirement: The concentration of the test chemical in the test containers should be determined at the beginning and end of the test. Modification: Based on previous testing, there was potential for the test material concentrations to decline rapidly in the exposure. Therefore, test solution samples were collected daily for analysis to better define the exposure concentrations. Additionally, the study results are presented on initial exposure concentrations and the time-weighted average (T.W.A.). The T.W.A. concentrations were calculated by averaging the measurements for the time interval with the measurement from the previous interval(s).

Additional modifications of the guideline requirements:

Requirement: Algal stock cultures should be shaken twice daily by hand. Test cultures should be oscillated on a rotary shaker at 100 rpm. Modification: Stock cultures were maintained, as were the test cultures, on a rotary shaker at 100 rpm.

Requirement: No chelating agents are to be included in the nutrient medium used for test solution preparation. Modification: Based on a recent request by U.S. EPA, Na_2EDTA . $2H_2O$ was added to the test medium at a concentration of 0.38 mg/l to promote algal growth.

Requirement: Designation of the test organism strain. Modification: The strain is not available for the test species *Selenastrum capricornutum*.

RESULTS

Nominal concentrations of 10, 5.0, 2.5, 1.3, 0.63 and 0.33 mg A.I./l were selected for the definitive test.

Initial measured concentrations used in this test were: 7.2, 2.9, 2.1, 1.2, 0.63 and 0.33 mg A.I./l. (85% of nominal on average).

Time-weighted average concentrations were: 2.2, 0.95, 0.64, 0.34, 0.18 and 0.086 mg A.I./l

All results are expressed in mg A.I./l (based on HPLC-UV analysis):

Results based on initial measured concentrations with 95% confidence limits:

24-hour EC10 value, 1.5 mg A.I./l (-3.2 – 2.3 mg A.I./l)

24-hour EC50 value, 2.3 mg A.I./l (0.46 – 3.5 mg A.I./l)

24-hour EC90 value, 3.2 mg A.I./l (2.4 – 6.7 mg A.I./l)

48-hour EC10 value, 1.7 mg A.I./l (0.20 – 4.5 mg A.I./l)

48-hour EC50 value, 3.5 mg A.I./l (1.0 – 12 mg A.I./l)

48-hour EC90 value, 7.1 mg A.I./l (2.6 – 41 mg A.I./l)

72-hour EC10 value, 1.7 mg A.I./l (0.71 – 4.2 mg A.I./l)

72-hour EC50 value, 3.6 mg A.I./l (1.5 – 9.3 mg A.I./l)

72-hour EC90 value, 7.5 mg A.I./l (3.2 – 22 mg A.I./l)

96-hour EC10 value, 0.98 mg A.I./l (-1.2 - 3.0 mg A.I./l)

96-hour EC50 value, 3.9 mg A.I./l (1.9 – 6.1 mg A.I./l)

96-hour EC90 value, 6.9 mg A.I./l (4.8 – 9.3 mg A.I./l)

Results based on time-weighted average concentration with 95% confidence limits:

24-hour EC10 value, 1.0 mg A.I./l (-2.4 – 1.7 mg A.I./l)

24-hour EC50 value, 1.7 mg A.I./l (0.27 – 2.5 mg A.I./l)

24-hour EC90 value, 2.3 mg A.I./l (1.7 – 5.0 mg A.I./l)

48-hour EC10 value, 0.85 mg A.I./l (0.20 – 2.1 mg A.I./l)

48-hour EC50 value, 1.7 mg A.I./l (0.59 – 4.8 mg A.I./l)

48-hour EC90 value 3.4 mg A.I./l (1.4 – 14 mg A.I./l)

72-hour EC10 value, 0.65 mg A.I./l (0.26 – 1.6 mg A.I./l)

72-hour EC50 value, 1.4 mg A.I./l (0.58 – 3.6 mg A.I./l)

72-hour EC90 value, 3.0 mg A.I./l (1.2 – 8.6 mg A.I./l)

96-hour EC10 value, 0.30 mg A.I./l (-0.31 – 0.87 mg A.I./l)

96-hour EC50 value, 1.2 mg A.I./l (0.63 – 1.8 mg A.I./l)

96-hour EC90 value, 2.1 mg A.I./l (1.5 – 2.8 mg A.I./l)

The 96-hour NOEC value based on initial measured concentration was 2.1 mg A.I./l and 0.64 mg A.I/l based on time weighted average concentration.

All measured concentrations decreased over time. At test termination, only the two highest test concentrations contained measurable amounts of test material (detection limit = 0.025 mg A.I./l). Test vessels containing 1.0 mg A.I./l solutions, but without algae, were analysed at 0, 2, 4, 8, 12, 24, 48, 72 and 96 hours to determine the loss of 2,6-DTBP in the absence of algae. After 4 hours, measured concentrations in these solutions declined faster than those in the test solutions containing algae. The decline in concentrations of 2,6-DTBP was apparently not caused by the algae in those solutions.

The HPLC-RAM analysis of the test solutions was performed to confirm the HPLC-UV analytical results and identify any radiolabelled degradates of 2,6-DTBP. The resultant measured concentrations were similar to those determined by HPLC-UV analyses. Although the concentrations of 2,6-DTBP decreased over time, no radiolabelled degradates were detected. It is inferred that the degradates of 2,6-DTBP in water were volatile and were released to the headspace above the test solution. Because all of the ¹⁴C in the test solutions occurred as 2,6-DTBP, total ¹⁴C measurement using LSC were employed for analysis of the 48, 72 and 96-hour samples to maximise the analytical sensitivity.

The disappearance half-life of 2,6-DTBP in each test concentration was calculated from the slope of the log concentration vs. time regression. The half-life of the test material ranged from 14 to 17 hours. The half-life of the un-inoculated 1.0 mg A.I./l test solutions was estimated to be 8.4 hours.

Cell density results:

Cell density for each flask at 24, 48, 72 and 96 hours was respectively:

7.2 mg A.I./l initial measured concentration: 2, 2, 1, 2 (x 10⁴ cells/ml)

2.9 mg A.I./l initial measured concentration: 1, 7, 6, 9 (x 10⁴ cells/ml)

2.1 mg A.I./l initial measured concentration: 4, 14, 15, 18 (x 10⁴ cells/ml)

1.2 mg A.I./l initial measured concentration: 4, 13, 16, 18 (x 10⁴ cells/ml)

0.63 mg A.I./l initial measured concentration: 3, 13, 19, 19 (x 10⁴ cells/ml)

0.33 mg A.I./l initial measured concentration: 5, 14, 18, 20 (x 10⁴ cells/ml)

Solvent control: 6, 16, 14, 21 (x 10⁴ cells/ml)

Control: 6, 16, 14, 19 (x 10⁴ cells/ml)

Pooled control: 6, 16, 14, 20 (x 10⁴ cells/ml)

Reference culture: not performed at the 24-hour interval, 10, 29, 83 (x 10⁴ cells/ml)

Control response was satisfactory.

Control culture averaged 19×10^4 cells/ml at 96 hours and solvent control cultures averaged 21×10^4 cells/ml. Cell densities in the reference cultures averaged 83×10^4 cells/ml at 96 hours. The lower densities observed in the controls and solvent controls presumably reflect carbon dioxide limitation in the sealed flask. Cell densities in the cultures exposed to 2,6-DTBP ranged from 2 to 20×10^4 cells/ml at 96 hours and generally followed the concentration gradient. Cell growth was not completely inhibited in any of the concentrations tested.

CONCLUSIONS

Based on time-weighted average concentrations the 96-hour EC50 value was 1.2 mg A.I./l and the 96-hour NOEC value 0.64 mg A.I./l.

DATA QUALITY

GLP study

Method follows EPA/TSCA guideline.

Information taken from the test report.

REFERENCES

2,6-Di-tert-butylphenol (DTBP) – Toxicity to the Freshwater Green Alga, *Selenastrum capricornutum* (final report) with attachments and covering letter dated 11.07.91

OTHER

Study Ref.: 91-7-3822, Springborn Laboratories, Inc., Environmental Sciences Division, 790 Main Street, Wareham, Massachusetts 02571, USA.

11.3 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

Purity of the test substance: information not available.

METHOD

Toxicity to Aquatic Plants

Test type: Data not available

GLP: (Y)

Species/strain: Algae (Selenastrum capricornutum)

Year study performed: 1994

RESULTS

Two attempts were made to test algal toxicity in response to the U.S. EPA proposed Test Rule for 2,6-DTBP. Both studies were found to be unacceptable, because stable test concentrations could not be maintained during the test period. EPA's Environmental Effects Branch finally concluded that a meaningful DTBP toxicity value for algae cannot be attained using the current or modified OPPT test guideline and EPA decided that no additional algal testing would be required.

CONCLUSIONS

No toxicity value for 2,6-DTBP was obtained in this test.

DATA QUALITY

GLP study

Information taken from the OECD/SIDS data set.

Purity of the test substance: information not available.

REFERENCES

Secondary reference: OECD/SIDS. Screening Information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref.: IRPTC Data Profile

11.4 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2,6-Di-*tert*-butylphenol

CAS No. 128-39-2

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.65 mg/l

ChV (96hr) 0.27 mg/l

Remarks: log Kow used 4.48 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.65 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

The test substance used in this test was 98.4% pure.

The test substance was identified on the basis of the code number. The characterisation and analysis were not part of the study.

METHOD

OECD Guideline No. 202 Acute Toxicity to Daphnia magna (24 hr)

Test type: Static

GLP: Data not available

Year study performed: 1987

Species: Daphnia magna Straus 1820

Analytical procedures: Data not available

Test details: The test was conducted under static conditions, in beakers covered with watch glasses.

Statistical methods: The EC50 values were calculated according to J. Berkson, Jasa 48 (1953), 569-599. EC50 was graphically determined on gausso-logarithmic probability paper.

RESULTS

Nominal concentrations: 0.58, 1.00, 1.80, 3.20 and 5.30 mg/l

All results are expressed in mg/l and are based on the nominal concentrations.

Calculated EC50 value with 95% confidence limits:

EC50 (24 hr) 1.7 mg/l (1.5-2.0 mg/l)

Graphically determined EC0, EC50 and EC100 values:

EC0 (24 hr) 0.58 mg/l EC50 (24 hr) 1.8 mg/l EC100 (24 hr) 5.8 mg/l

Number immobilised as compared to the number exposed to nominal concentrations:

0.58 mg/l: 0/20 (0%) 1.0 mg/l: 1/20 (5%) 1.80 mg/l: 14/20 (70%) 3.20 mg/l: 15/20 (75%) 5.80 mg/l: 20/20 (100%)

Controls: immobilisation blank 0/20 (0%)

immobilisation vehicle 0/20 (0%)

Control response was satisfactory.

CONCLUSIONS

EC50 (24 hr) value for 2,6-Di-tert-butylphenol was found to be 1.7 mg/l.

DATA QUALITY

Method follows OECD guideline.

Information taken from the test report.

GLP study: data not available

REFERENCES

Report on the test for acute toxicity of TK 12891 to *Daphnia magna*, Ciba-Geigy Ltd, AD-Division, CH-4002, Basel, Switzerland.

OECD/SIDS. Screening Information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref.: 87 40 57 – Unpublished Report

12.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

METHOD

Acute Toxicity to *Daphnia magna* (24 hr)

Official Journal of the European Communities L251C-02, 19.9.1984

Test type: Static

GLP: Data not available

Year study performed: 1990

Species: Daphnia magna Straus 1820

Analytical procedures: Data not available.

Remarks: Small parts of test substance were floating at the surface of the test vessels at 10 and 18 mg/l after 24 hours.

Test details: The test was conducted under static conditions, in beakers covered with watch glasses.

Statistical methods: The EC50 values were calculated according to the maximum likelihood method probit model (Mc Cullagh, P., Nelder, J.A., 1983: Generalised linear models, Chapman & Hall, London).

EC values were graphically determined on gausso-logarithmic probability paper.

There were no deviations to protocol.

RESULTS

Nominal concentrations: 0.32, 0.58, 1.0, 1.8, 3.2, 5.8, 10 and 18 mg/l

All results are expressed in mg/l and are based on nominal concentrations.

Calculated EC50 value with 95% confidence limits:

EC50 (24hr) 5.5 mg/l (4.5-6.6 mg/l)

Graphically determined EC0, EC50 and EC100 values:

EC0 (24hr) 1.8 mg/l EC50 (24hr) 5.0 mg/l EC100 (24hr) 18 mg/l

Number immobilised as compared to the number exposed to nominal concentrations:

0.32 mg/l: 0/20 (0%) 0.58 mg/l: 0/20 (0%) 0/20 (0%) 1.0 mg/l: 1.8 mg/l: 0/20 (0%) 3.2 mg/l: 2/20 (10%) 5.8 mg/l: 15/20 (75%) 10 mg/l: 15/20 (75%) 18 mg/l: 20/20 (100%)

Controls: immobilisation blank 0/20 (0%)

immobilisation vehicle 0/20 (0%)

Control response was satisfactory.

CONCLUSIONS

Calculated EC50 (24hr) value for 2,6-Di-tert-butylphenol was found to be 5.5 mg/l.

DATA QUALITY

Information taken from the test report.

Purity of the test substance: information not available.

GLP study: no data available.

REFERENCES

Report on the acute toxicity test of IRGANOX 108 to *Daphnia (Daphnia magna* Straus 1820), Ciba-Geigy Ltd, AD-Division, CH-4002, Basel, Switzerland.

OTHER

Study Ref.: 094580

12.3 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 1.1 mg/l ChV (21day) 0.098 mg/l

Remark: log Kow used 4.48 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 1.1 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

12.4 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA) - ADDITION

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

The test substance used in this test was 99.88% pure.

METHOD

EPA/ OTS Standards

Test type: Flow through Test

GLP: Y

Year study performed: 1987

Species: Daphnia magna (Water flea)

Analytical procedures: Data not available

Exposure Period: 24-48 hours

Dose/ Concentration: 0.076 – 0.59 mg/l

RESULTS

EC50 (24 hr) (behavioral) >0.59 mg/l

EC50 (48 hr) (behavioral) 0.45 mg/l

NOEC (behavioral) 0.076 mg/l

CONCLUSIONS

EC50 (48 hr) value for 2,6-Di-tert-butylphenol based on behavioral changes was found to be 0.45 mg/l.

DATA QUALITY

Method follows EPA/ OTS standards.

GLP compliant.

REFERENCES

Ethyl Corp. Unpublished report – Ethyl Corporation, 89-05-2948 (1989).

OECD/SIDS. Screening Information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

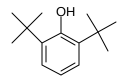
12.4 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., SCUD) - ADDITION

TEST SUBSTANCE

2,6-Di-tert-butylphenol

CAS No. 128-39-2

Purity of Test Substance: 99.88%



METHOD

Test type: Flow through Test

GLP: Y

Year study performed: 1988

Species: Gammarus fasciatus (Scud)

Analytical procedures: Data not available

Exposure Period: 24-96 hours

Dose/Concentration: 0.6 - 1.0 mg/l and additional dose of 0.38 mg/l was run for 96 hours.

Solvent: Acetone

RESULTS

LC50 (24 hr) 1.0 mg/l

LC50 (48 hr) 0.80 mg/l

LC50 (72 hr) 0.70 mg/l

LC50 (96 hr) 0.60 mg/l

NOEL (96 hr) 0.38 mg/l

CONCLUSIONS

LC50 (96 hr) value for 2,6-Di-tert-butylphenol was found to be 0.60 mg/l.

DATA QUALITY

GLP compliant.

REFERENCES

Ethyl Corp. Unpublished report – Ethyl Corporation, 88-12-2881 (1988).

OECD/SIDS. Screening Information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

No information is provided concerning the degree of purity of the material or the impurities used in the test.

METHOD

Interstate Commerce Commission, Tariff No. 8 Section 73.343 Acute Oral Toxicity Test

GLP: (N)

Year study performed: 1955

Species: Rat

Sex: Female

No of animals per sex per dose: 10

Vehicle: Substance as such

Route of administration: Oral gavage

Remarks: Ten rats were given 0.2 ml of the test substance maintained at 38°C, per kg of body weight, by means of a blunt needle which traversed the oesophagus.

RESULTS

All animals survived and showed no signs of illness, other than slight losses in body weight gain.

LD50 7ml/kg (6286 mg/kg) approximate

CONCLUSIONS

2,6 di-tertiary butyl phenol was far less toxic then the di-iso-propyl phenol when administered orally to rats.

Information in this test report may not be acceptable due to deficiency of data element. However, this test will be retained in the file for valuable comparison and overall evaluation of toxicity of the test material.

DATA QUALITY

Very old study with no GLP.

REFERENCES

The Immediate Toxicity of 2,6-Diisopropylphenol and 2,6-Di-tertiary Butyl Phenol, in Relation to the Regulations of the Interstate Commerce Commission for the Transport of Chemicals. Kettering Laboratory, Dept of Preventative Medicine and Industrial Health, College of Medicine, University of Cincinnati, Cincinnati, Ohio, USA.

OTHER

Ref: 42096 B3-8

13.2 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guidelines No. 401

GLP: (Y)

Year study performed: 1991

Species: Sprague-Dawley strain rat

Sex: Males and females

No of animals per sex per dose: 5000 mg/kg, Five males/five females

Vehicle: The test material was prepared, as a solution/suspension at the appropriate concentration in Arachis oil B.P. Homogeneity was assured by the use of a Silverson Homogeniser.

Route of administration: Oral, gavage

Remarks: At the start of the main study the animals were approximately five to eight weeks old. Following the range finding study, a group of ten fasted animals were given a single oral dose of the test material, as a solution/suspension in Arachis oil B. P. at a dose level of 5000 mg/kg. Animals were observed for fourteen days.

RESULTS

LD50 value of the test material was found to be greater then 5000 mg/kg body weight. Two animals (one male and one female) were killed one day after treatment. Common signs of systemic toxicity noted were hunched posture, lethargy, ataxia and occasional body tremors with additional signs of laboured respiration, loss of righting reflex and ptosis. Surviving animals appeared normal 3 days after treatment and showed expected gain in body weight during the study.

Abnormalities noted at necropsy of animals killed during the study were abnormally red lungs, patchy pallor of the liver, dark kidneys, hemorrhage of the gastic mucosa and hemorrhage of the small and large intestines. No abnormalities were noted at necropsy of animals killed at the end of the study.

CONCLUSIONS

The test substance has not been classified in this Acute Oral Toxicity Test.

LD50: >5000 mg/kg

DATA QUALITY

Performed to OECD guidelines and to GLP.

REFERENCES

DTBP-Range Finding Acute Oral Toxicity Test in the Rat, Safepharm Laboratories Ltd, P.O. Box No. 45, Derby, DE1 2BT, U.K.

OTHER

Study Ref.: 47/1608

13.3 ACUTE TOXICITY INHALATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Interstate Commerce Commission, Tariff No. 8 Section 73.343 Acute Inhalation Toxicity Test

GLP: (N)

Year study performed: 1955

Species: Rat

Sex: Female

No of animals per sex per dose: 2 females

Vehicle: Air was bubbled through the test substance, maintained at 100°C in an absorption tower, and passed through the 5.82 l chamber at a rate of approximately 515 ml/min. The amount of compound that was carried out of the tower by the flowing air was estimated by weighing the absorption tower containing the compound before starting and after terminating the exposure.

Route of administration: Inhalation - vapor

Remarks: Only one experiment was performed. The animals were exposed to the test material for 3 hours.

RESULTS

No more than traces of the test substance were carried into the chamber by the flowing air and these failed to cause any signs of illness in the rats.

CONCLUSIONS

The vapor pressure of the test substance was not sufficiently high to provide a toxic or lethal concentration within the atmosphere of the respiratory chamber under the conditions (duration of exposure) of the experiment.

Information in this test report may not be acceptable due to deficiency of data element. However, this test is included for valuable comparison and overall evaluation of toxicity of the test material.

DATA QUALITY

Not conducted to GLP.

REFERENCES

The Immediate Toxicity of 2,6-Diisopropylphenol and 2,6-Di-tertiary Butyl Phenol, in Relation to the Regulations of the Interstate Commerce Commission for the Transport of Chemicals. Kettering Laboratory in the Department of Preventative Medicine & Industrial Health, College of Medicine, University of Cincinnati, Cincinnati, Ohio, U.S.A.

OTHER

Study Ref: 42096 B3-8

13.4 ACUTE TOXICITY DERMAL

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

No data available

GLP: No data

Year study performed: No data

Species/strain: rat

Route of administration: dermal

RESULTS

Dermal LD50 for rats estimated as greater than 1000 mg/kg and less then 32000 mg/kg.

The given LD50 is according to information submitted under TSCA Section 8(d) by Ethyl Corporation.

CONCLUSIONS

LD50 > 1000 mg/kg and < 32000 mg/kg

DATA QUALITY

Data taken from a published paper

REFERENCES

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

13.5 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guidelines No. 404

GLP: (Y)

Year study performed: 1991

Species: New Zealand White rabbits

Sex: Males and females

No of animals per sex per dose: One male and two females

Vehicle: 0.5 g of the test material was ground and moistened with 0.5 ml of distilled water immediately before application.

Remarks: A group of three rabbits were given a single 4-hour, semi-occluded application of the test material to the intact skin. The animals were observed for fourteen days after the day of the dosing.

RESULTS

One animal was killed for humane reasons nine days after patch removal due to illness.

Very slight to well-defined erythema was noted at all treated skin sites one hour after patch removal, with well-defined erythema at all treated skin sites after the 24, 48 and 72-hour observation. Other adverse skin reaction noted at one treated skin site at that time were hemorrhage of dermal capillaries, light brown discoloration of the epidermis, thickening of the skin and loss of skin elasticity. Crust formation was noted at the 7-day observation. Hardened light brown-colored scabs and thickening of the skin were noted at two treated skin sites at the 7-day observation. These reactions prevented accurate evaluation of erythema and edema. Crust formation and/or reduced re-growth of fur were noted at two treated skin sites at the 14-day observation. Very slight to moderate edema was noted at all treated skin sites one and 24 hours after patch removal with very slight to slight edema at the 48 and 72-hour observations.

CONCLUSIONS

The test material produced a primary irritation index of 3.8 and was classified as a moderate irritant to rabbit skin according to the Draize classification scheme. The test material produced positive criteria in 3/3 rabbits according to the EEC labelling regulations and was classified as irritant to rabbit skin. The symbol Xi and the risk phrase R38 "Irritating to skin" are therefore required.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

DTBP-Acute Dermal Irritation Test in the Rabbit, Safepharm Laboratories Ltd, P.O. Box No. 45, Derby, DE1 2BT, U.K.

OTHER

Study Ref: 47/1609

14.1 GENETIC TOXICOLOGY IN VITRO

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

Guidelines followed: Protocol-SOP ETTOX 029 with modifications of the method as described by Williams (1978)

Type: Rat hepatocyte primary culture/DNA repair test

GLP: (Y)

Year: 1984

Species: Fischer-344 rats

Sex: Male

Weight at initiation: 150-300 grams (adult rats)

The rats were anaesthetised with sodium numbutal by intraperitoneal injection. Following anaesthetisation, the livers were perfused and removed. The livers were excised and the hepatocytes harvested.

Freshly isolated rat hepatocytes were treated with 20 μ l of test substance at 0.05, 0.1, 0.5, 1, 5, 10, 50, 100, 500 and 1000 μ g/well in 2 ml of media. The hepatocytes were then fixed onto microscope coverslides, stained, dipped and developed.

Unscheduled DNA repair synthesis was quantified by a net increase of black silver grains in the nucleus for 25 cell/slide. This value was determined by taking a nuclear count and the average of three adjacent cytoplasmic counts.

RESULTS

Cytotoxicity was produced at concentrations of 50, 100, 500 and 1000 μ g/well. The highest dose scored was therefore 10 μ g/well. The test substance did not cause an increase in mean net nuclear counts over DMSO (negative control) at any dose level counted. A Chi square analysis was performed to compare treated cells to untreated controls. The number of cells with net nuclear grain counts greater than zero of all scored dose levels of test substance was not statistically significantly increased.

CONCLUSIONS

The test substance was not genetically active in this in vivo assay.

DATA QUALITY

Study conducted to GLP and guidelines.

REFERENCES

Genetic toxicology- Rat hepatocyte primary culture/DNA repair test Report number HPC 022 - #068

Ethyl Corporation 8000 GSRI Avenue, Baton Rouge, Louisiana 70808, U.S.A.

15.1 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

Guideline followed: Ames test

METHOD

Type: Plate incorporation assay

System of testing: Bacterial

S.O.P. ETTOX-027

GLP: (Y)

Year study performed: 1984

Species/strain: Salmonella typhimurium, TA 1535, TA 1537, TA 98 and TA 100.

Metabolic activation: *Salmonella typhimurium* strains were treated in the presence and absence of metabolic activation Aroclor 1254 induced in rat liver S9 fraction with the test substance in acetone.

Concentrations tested: $0, 2.5 \times 10^{-2}, 5 \times 10^{-3}, 2.5 \times 10^{-3}, 5 \times 10^{-4}$ and 2.5×10^{-4} mg/plate

Remarks: Strains were treated in the presence and absence of metabolic activation with the test substance dissolved in acetone. Each dose was treated in triplicate. An untreated control, solvent control and positive control were treated concurrently.

The solvent control for TA 98 with activation was unacceptably below the spontaneous rate and the historical solvent control rate. There was no significant increase of the treated in mutant colonies over the spontaneous rate. Therefore, the spontaneous rate was used as the negative control in TA 98 with activation.

RESULTS

 2.5×10^{-2} mg/plate demonstrated toxicity in strains TA 1535, TA 1537 and TA 100 with and without metabolic activation.

Negative: The test substance did not induce a dose-related increase in mutant colonies over acetone in any strain in the absence or presence of metabolic activation.

CONCLUSIONS

The test material did not cause a dose-related increase in mutant colonies and is not genetically active in the Ames *Salmonella typhimurium* assay.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

Genetic Toxicity; Salmonella/Microsomal Assay

OTHER

Study Ref: Ames test 086-068 /Ethyl Corporation, 8000 GSRI Avenue, Baton Rouge, Louisiana 70808, U.S.A.

15.2 GENETIC TOXICITY IN VITRO (GENE MUTATIONS)

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHODS

Guideline followed: Ames test Type: Reverse mutation assay

System of testing: Bacterial

GLP: No data available

Year study performed: 1985

Species/strain: Escherichia coli: WP2, WP2 uvrA

Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA 1538

Metabolic activation: Strains were treated in the presence and absence of metabolic activation by S-9 mix.

Concentrations tested: No data available

RESULTS

The results of the test were negative for mutagenic effects in the presence or absence of metabolic activation.

CONCLUSIONS

The test material is not genetically active in the Salmonella and Escherichia coli assay.

DATA QUALITY

Study conducted according to guidelines but no data to show if to GLP.

REFERENCES

Primary reference:

Dean et al. Mutation Research, 153(1-2), 57-77, (1985)

Secondary reference: OECD/SIDS. Screening Information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1984)

OTHER

Study ref: IRPTC data profile

15.3 GENETIC TOXICITY IN VITRO (CHROMOSOMAL ABERRATIONS)

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guideline No 473; EEC Directive 84/449, B 10

Type: Chromosome aberration assay System of testing: Non bacterial

GLP: (Y)

Year study performed: 1992

Species/strain: Chinese hamster V79 cells

Metabolic activation: S-9 mix (with and without)

Exposure period: 18 to 28 hours

Concentrations tested:

Dose/concentration without S -9 mix: 18 hr - 0.3, 3.0, 6.0, 10.0 μ g/ml 28 hr - 6.0 μ g/ml

Dose/concentration with S-9 mix: $18 \text{ hr} - 3.0, 30.0, 50.0 \,\mu\text{g/ml}$

 $28 \text{ hr} - 50.0 \ \mu\text{g/ml}$

RESULTS

No mutagenic effects were obtained with or without metabolic activation. The test did not produce chromosomal aberrations.

The test substance negatively affected the plating efficiency of the cells at concentrations higher than $3.0\,\mu\text{g/ml}$ (without S-9 mix) and $30.0\,\mu\text{g/ml}$ (with S-9 mix).

CONCLUSIONS

The test substance is not considered to be mutagenic.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

Primary reference:

Sandoz Chemical Ltd. Muttenz, Switzerland. Unpublished CCR Report- Sandoz, 243628, (1992)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

15.4 GENETIC TOXICITY IN VITRO (GENE MUTATION ASSAY)

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guideline No 476; 40 CFR, Ch. I part 798

Type: Mammalian cell gene mutation assay

System of testing: Non bacterial

GLP: (Y)

Year study performed: 1991

Species/strain: Chinese hamster V79 cells

Metabolic activation: S-9 mix (with and without)

Concentrations tested:

Dose/concentration without S -9 mix: 0.3, 1.0, 2.0, 4.0, 6.0 and 8.0 μ g/ml Dose/concentration with S-9 mix: 3.0, 10.0, 20.0, 30.0, 40.0 and 50.0 μ g/ml

RESULTS

Up to the highest concentrations of the test substance no relevant increase in mutant colony numbers was obtained in two independent separate experiments. There was no mutagenic effect observed with or without metabolic activation.

CONCLUSIONS

The test substance is not considered to be mutagenic.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

Primary reference:

Sandoz Chemical Ltd. Muttenz, Switzerland. Unpublished CCR Report- Sandoz, 243617, (1991)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

16.1 REPEATED DOSE TOXICITY 28 DAYS ORAL

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

EEC Directive 84/449/EEC and OECD Guideline No. 407

GLP: No data available

Year study performed: 1992

Species/strain: Rat, Wistar

Sex: Males and females

No of animals per sex per dose: 5 males and 5 females

Route of administration: Oral, gavage

Exposure Period: 28 days

Dose/Concentration: 0, 15, 100 and 600 mg/kg/day

RESULTS

NOAEL 15 mg/kg/day

At the doses of 100 and 600 mg/kg/day, there were decreased levels of serum urea in females only.

Increased relative liver weights were found at 100 mg/kg/day dose level in males only.

At macroscopic examination enlarged caecum was noted in 2 out of 5 males in the dose group of 100 mg/kg/day.

An increased serum total protein level was found in males and females, and increased serum albumin level was found in males at the dose group of 600 mg/kg/day.

Decreased inorganic phosphate and increased calcium levels were noted in the serum of females from the 600 mg/kg/day dose groups.

In the dose group of 600 mg/kg/day enlarged caecum was found in 4/5 males and 5/5 females. In the same dose group enlargement of liver and kidneys were noted. At 600 mg/kg/day increased liver weights were noted in males and females, and increased kidney weights in males only.

At microscopic examination, in the 600 mg/kg/day dose group there was a slight increase of hepatocellular hypertrophy in the centrilobular area in males and females, and eosinophilic inclusions in the renal cortex of males only.

CONCLUSIONS

Concentration of the test substance at which no toxic effects were observed was 15 mg/kg/day under the test conditions.

DATA QUALITY

Study conducted according to OECD guidelines but no data to show if to GLP.

REFERENCES

Primary reference:

Sandoz Chemical Ltd, Muttenz, Switzerland. Unpublished CCR Report- Sandoz, 304435, (1992)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

17.1 TOXICITY TO REPRODUCTION

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guideline No. 421 "Preliminary Reproduction/ Developmental Toxicity Screening Test"

GLP: (Y)

Year study performed: 1992

Species/strain: Rats, Wistar

Route of administration: Oral gavage

Dose/concentrations levels: 0, 30, 150 and 750 mg/kg/day

Sex: Males and females

Control group and treatment: 10 males, 10 females per group

Frequency of treatment: 7 days/week

Duration of test: From 2 weeks prior to mating until 4 days post partum (study termination)

Pre-mating exposure period for males: 2 weeks

Pre-mating exposure period for females: 2 weeks

Animals were dosed throughout the pre-mating and mating period (up to 13 days) for all animals. Males received the test substance for a further 43 days and females up to Day 3 post-partum.

RESULTS

Marginally reduced body weight gain was observed in male rats from the dose group at 750 mg/kg/day, in spite of increased food consumption. Slight reduction of body weight gains was observed in females of 750 mg/kg/day. Reduced food consumption was observed in these females.

The mutual reproductive parameters (pre-coital time, percentage mating, fertility index and conception rate) indicated no test substance related effects in any dose group. An increased breeding loss/ reduced viability index was observed for the females of the higher dose group (750 mg/kg/day).

Neither macroscopic examination of the parent animals, nor microscopic examination or mean organ weight and organ/body weight ratios of testes and ovaries, gave any indication of test substance related effects.

Based on these results, the only observed effect was the appearance of severe toxic symptoms in the higher dose group (750 mg/kg/day). With respect to the reproductive and developmental parameters, no effects were noted up to and including 150 mg/kg/day.

NOAEL – Systemic and reproductive = 150 mg/kg/day

The summary information on the reproductive study for CAS No. 128-39.2 was apparently transcribed from the original report and published in the OECD SIDS. It appears that the transcription was not without possible error(s). The NOAEL provided in the SIDS is stated to be 750 mg/kg/day. However, examination of the summary results suggests this value should be 150 mg/kg/day, based on clinical symptoms, reduced food consumption and reduced body weight gain in the dams at the 750 mg/kg/day dose level.

The NOAEL for developmental toxicity is stated to be 750 mg/kg/day. However, at 750 mg/kg/day a reduction in pup body weight gain was reported in the summary, but not further described as to duration or extent. In addition, the pup viability index (either measured on lactation day 4 or 21) was apparently reduced. For these reasons, and without further study details, it is probably more appropriate to consider this NOAEL as being 150 mg/kg/day and not 750 mg/kg/day as stated in the summary.

CONCLUSIONS

The test substance is not toxic for reproduction in rats.

DATA QUALITY

The study was conducted to GLP and OECD guidelines

REFERENCES

Primary reference:

Sandoz Chemical Ltd. Muttenz, Switzerland. Unpublished CCR Report- Sandoz, 321794, (1992)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

18.1 DEVELOPMENTAL TOXICITY/TERATOGENICITY

TEST SUBSTANCE

2,6-Di-tert butylphenol

CAS No. 128-39-2

METHOD

OECD Guideline No. 421 "Preliminary Reproduction/ Developmental Toxicity Test"

GLP: (Y)

Year study performed: 1992

Species/strain: Rat, Wistar

Route of administration: Oral, gavage

Dose/concentration: 0, 30, 150 and 750 mg/kg/day were tested.

Sex: Males and females

Control group and treatment: 10 males, 10 females per group

Frequency of treatment: 7 days/ week

Duration of test: Treatment from 2 weeks prior to mating until 4 days post-partum (study termination)

Organs examined at necropsy (macroscopic and microscopic): Data not available

RESULTS

NOAEL maternal toxicity 750 mg/kg/day

NOAEL developmental toxicity >750 mg/kg/day

Maternal data:

The only observed effect was the appearance of severe toxic symptoms in the higher dose group (750 mg/kg/day). With respect to the reproductive and developmental parameters, no effects were noted up to and including 750 mg/kg/day.

Neither macroscopic examination of the parent animals, nor microscopic examination or mean organ weight and organ/body weight ratios of testes and ovaries, gave any indication of test substance related effects.

Fetal data:

At 750 mg/kg/day maternal exposure dose level the body weight gain of pups was reduced. Other than reduced body weight gain, no test substance related effects on the pups were noted in any dose group.

Remarks:

The summary information on the reproductive study for CAS No. 128-39.2 was apparently transcribed from the original report and published in the OECD SIDS. It appears that the transcription was not without possible error(s). The NOAEL provided in the SIDS is stated to be 750 mg/kg/day. However, examination of the summary results suggests this value should be 150 mg/kg/day, based on clinical symptoms, reduced food consumption and reduced body weight gain in the dams at the 750 mg/kg/day dose level.

The NOAEL for developmental toxicity is stated to be 750 mg/kg/day. However, at 750 mg/kg/day a reduction in pup body weight gain was reported in the summary, but not further described as to duration or extent. In addition, the pup viability index (either measured on lactation day 4 or 21) was apparently reduced. For these reasons, and without further study details, it is probably more appropriate to consider this NOAEL as being 150 mg/kg/day and not 750 mg/kg/day as stated in the summary.

CONCLUSIONS

The test substance did not cause any reproductive/developmental effects in Wistar rats.

DATA QUALITY

Study conducted to GLP and OECD guidelines

REFERENCES

Primary reference:

Sandoz Chemical Ltd. Muttenz, Switzerland. Unpublished CCR Report-Sandoz, 321794, (1992)

Secondary reference:

OECD/SIDS. Screening information Data Set (SIDS) of OECD High Production Volume Chemicals Programme, (1994)

OTHER

Study Ref: IRPTC Data Profile

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,4-Di-*tert*-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1932.

RESULTS

Melting point: 26℃.

CONCLUSIONS

The test substance has a melting point of 26°C.

DATA QUALITY

Not a GLP study

Information taken from the results of a literature search covering appropriate databases.

Purity of the substance / decomposition: information not available.

REFERENCES

Dow Chem. Co., US 1972599 1932, Handbook Data

TEST SUBSTANCE

2,4-Di-tert-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Boiling point: 310.95°C.

CONCLUSIONS

The test substance has a calculated boiling point of 310.95°C.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30.

OTHER

Adapted Stein & Brown method used.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2,4-Di-*tert*-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 8.41 x 10⁵ mm Hg @ 25℃ (0.011 Pa).

CONCLUSIONS

The test substance has a calculated vapor pressure of $8.41 \times 10^{-5} \text{mm Hg} \ @ 25^{\circ}\text{C}$.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30

OTHER

Modified Grain method used.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,4-Di-tert-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 631

CONCLUSIONS

The test substance has a log Kow of 6.31.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,4-Di-*tert*-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation using a calculated log Kow value of 6.31.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 0.4441 mg/l @ 25°C.

CONCLUSIONS

The test substance has a calculated water solubility of 0.4441 mg/l.

DATA QUALITY

Calculation.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

The log Kow value of 6.31 calculated by the programme KOWWIN (see 4.1) was used in the calculation of water solubility.

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,4-di-tert-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $51.4 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.50 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.50 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,4-di-tert-pentylphenol

CAS No. 120-95-6

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

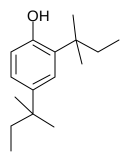
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,4-di-tert-pentylphenol

CAS No. 120-95-6



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass: 234

Data temperature (°C): 25

Water solubility (mg/l): 0.4441

Vapor pressure (Pa): 0.011

Log Kow: 6.31

Melting point (°C): 26

Environmental conditions: defaults used.

RESULTS

ENVIRONMENTAL COMPARTMENT	Percentage of test substance
Air	0.063
Soil	97.6
Water	0.054
Suspended sediment	0.068
Fish	0.0055
Sediment	2.17

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2,4-di-tert-pentylphenol

CAS No. 120-95-6

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in weeks

Ultimate biodegradation in weeks/months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2, 4-di-tert-Pentylphenol

CAS No. 120-95-6

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 95hours, 30days, 90 days

RESULTS

LC50 (96hr) 0.076 mg/l

ChV (30day) 0.011 mg/l

ChV (90day) 0.003 mg/l Remark: log Kow used 6.31 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) value for the test substance was found to be 0.076 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

11.1 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2, 4-di-tert-Pentylphenol

CAS No. 120-95-6

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.018 mg/l

ChV (96hr) 0.022 mg/l

Remark: log Kow used 6.31 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) value for the test substance was found to be 0.018 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2, 4-di-tert-Pentylphenol

CAS No. 120-95-6

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48hours, 21 days

RESULTS

LC50 (48hr) 0.22 mg/l

ChV (21day) 0.008 mg/l

Remark: log Kow used 6.31 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) value for the test substance was found to be 0.22 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,4-di-tert-pentylphenol

CAS No. 120-95-6

METHOD

Oral LD50 (Rats, mixed sex). Conducted according to a modification of the method by E.J. de Beer.

GLP: (N)

Year study performed: 1968

Species: Sprague-Dawley CD rats

Sex: Male and female

No of animals: 10 males, 10 females

Doses/concentrations: Animals were fed in increasing doses at increments of 0.1 fractional log intervals at four levels: 631, 794, 1000 and 1260 mg/kg/bodyweight.

Vehicle: No information given.

RESULTS

LD50: 920 mg/kg (confidence limits: 800 – 1060 mg/kg)

The substance is classified as mildly toxic by oral ingestion.

Survival time was 1 to 6 days with most deaths occurring in 3 to 4 days. Toxic symptoms included loss of appetite, increasing weakness, salivation, tremors and dyspnea. At autopsy there was liver discoloration, renal congestion and hemorrhagic areas in the lungs.

Animal/ Sex	Weight (g)	Dose (mg/kg)	Fate
Female	230	631	Survived
Male	245	631	Survived
Female	225	631	Survived
Male	260	631	Survived
Female	220	631	Survived
Male	250	794	Survived
Female	235	794	Died
Male	260	794	Survived
Female	215	794	Survived
Male	250	794	Survived
Female	230	1000	Survived
Male	255	1000	Died
Female	220	1000	Died
Male	245	1000	Survived
Female	235	1000	Died
Male	260	1260	Died
Female	240	1260	Died
Male	250	1260	Died
Female	230	1260	Died
Male	255	1260	Died

Remarks:

No deaths occurred at 631 mg/kg thus the test material could be classed as non toxic, however in the absence of any available information on bodyweight gains, clinical condition of the animals and

is likely the death at 794 mg/kg was treatment-related evidence of toxicity.

CONCLUSIONS

LD50: 920 mg/kg

DATA QUALITY

Study not performed to GLP.

REFERENCES

Monsanto Company, 800 N. Lindbergh Boulevard, St Louis, Missouri 63167, U.S.A. Testing performed by Younger Laboratories, Cliff Cave Road, St. Louis, Missouri 68189, U.S.A. project # Y-68-115

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: information not available.

GLP (Y/N): no.

Year study performed: 1960.

RESULTS

Melting point: 47℃.

CONCLUSIONS

The test substance has a melting point of 47°C.

DATA QUALITY

Not a GLP study.

Information taken from the results of a literature search covering appropriate databases.

Information on method, decomposition and purity not available.

REFERENCES

Ingold, J.Phys.Chem., 640 (1960), 1636, 1637. As quoted in a literature search covering appropriate databases.

OTHER

2.1 BOILING POINT

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): no.

Year study performed: 1993.

RESULTS

Boiling point: 275°C (atmospheric pressure assumed).

CONCLUSIONS

The test substance has a boiling point of 275° C.

DATA QUALITY

Not a GLP study.

Information taken from a technical datasheet.

Decomposition: information not available.

Purity of the test substance given as typically 95.0 % minimum.

REFERENCES

Schenectady International Inc. Technical Datasheet (11/93).

3.1 VAPOR PRESSURE

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: 2.07×10^5 mm Hg at 25 °C (0.0028 Pa).

CONCLUSIONS

The calculated vapor pressure is 2.07 $\times 10^{5}$ mm Hg at 25°C.

DATA QUALITY

Calculation.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBWIN v 1.30.

OTHER

Modified Grain Method used.

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 6.43.

CONCLUSIONS

The test substance has a calculated log Kow of 6.43.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: calculation method using a calculated log Kow of 6.43. GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 0.2479 mg/l at 25°C.

CONCLUSIONS

The calculated water solubility of the test substance is 0.2479 mg/l at 25°C.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

The log Kow value (7.17) calculated using the programme KOWWIN (see 4.1) has been used for the calculation of water solubility.

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $20.5 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 6.27 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 6.27 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

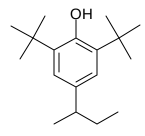
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

4-*sec*-Butyl-2,6-di-*tert*-butylphenol

CAS No. 17540-75-9



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:262Data temperature (°C):25Water solubility (mg/l):0.2479Vapor pressure (Pa):0.0028Log Kow:6.43Melting point (°C):47

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	0.0245
Soil	97.7
Water	0.041
suspended sediment	0.068
fish	0.0055
Sediment	2.17

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in weeks

Ultimate biodegradation in months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

4- sec- Butyl- 2, 6-di- tert-butylphenol

CAS No. 17540-75-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 95hours, 30days, 90 days

RESULTS

LC50 (96hr) 0.072 mg/l

ChV (30day) 0.010 mg/l

ChV (90day) 0.003 mg/l

Remarks: log Kow used 6.43 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) value for the test substance was found to be 0.072 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

11.1 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

4- sec- Butyl- 2, 6-di- tert-butylphenol

CAS No. 17540-75-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.016 mg/l

ChV (96hr) 0.020 mg/l

Remark: log Kow used 6.43 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) value for the test substance was found to be 0.016 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

4- sec- Butyl- 2, 6- di-tert-butylphenol

CAS No. 17540-75-9

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48hours, 21 days

RESULTS

LC50 (48hr) 0.22 mg/l

ChV (21day) 0.008 mg/l

Remark: log Kow used 6.43 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) value for the test substance was found to be 0.22 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v 0.99e

OTHER

Calculation performed for this HPV submission.

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

METHOD

Acute Oral LD50 Toxicity Study in Rats – according to the method of C.S. Weil, Biometrics 249 (1952)

GLP: (Y)

Year study performed: 1980

Species: Sprague-Dawley CD rats

Sex: Male and female

No of animals per sex per dose: 5 males, 5 females at each dose level

Doses/concentrations: 3.4, 4.7, 6.6 and 9.3 g/kg/bodyweight

Vehicle: Test articles in a dosing media prepared on a w/v basis.

RESULTS

Mortality summary (Number of deaths)									
Dosage	Hr 0-4	1	2	3	4	5	6	7-14	Totals
(g/kg)	M/F	M/F	M/F	M/F	M/F	M/F	M/F	M/F	Both
3.4	0/0	0/0	0/0	1/2	1/1	1/0	0/0	0/0	3/3
4.7	0/0	0/0	0/0	2/1	0/0	0/0	0/0	0/0	2/1
6.6	0/0	0/0	2/1	1/2	0/0	1/0	0/0	0/0	4/3
9.3	0/0	0/0	3/0	1/4	0/0	0/0	0/0	1/0	5/4

The LD50 with 95% confidence limits was estimated to be 4.8 (2.7 - 8.1) g/kg.

Clear evidence of toxicity was seen at all dose levels.

Clinical signs of toxicity included decreased motor activity, diarrhoea, piloerection, co-ordination loss and lethargy. Necropsy indicated irritation to the intestines.

Necropsy results:

Macroscopic examination at the study termination revealed bright red lungs, dark red mottled liver, intestinal irritation, nasal and ocular hemorrhage and diarrhoea in males and wet ventral surface in one female dosed at 3.4 g/kg.

In addition to these effects, in the test group dosed at 4.7 g/kg, slight hair loss was noted from ventral surface in one male and one female.

In the test group dosed at 6.6 g/kg and 9.3 g/kg necropsy examination revealed diarrhoea, lacrimation, wet ventral surface, stomach and intestinal irritation, stomach filed in food, hair loss from posterior ventral surface, lungs bright red mottled, small dark spots on thymus, and nasal and ocular hemorrhage in both males and females.

CONCLUSIONS

LD50 for 4-sec-butyl-2,6-tert-butyl phenol was found to be 4.8 g/kg.

DATA QUALITY

Study to GLP and guidelines

REFERENCES

Springborn Institute for Bioresearch Inc., Spencerville, Ohio 45887 Lab Study No.: 3076.16, Report date: 12/18/80.

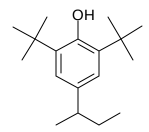
15.1 GENETIC TOXICITY IN VITRO (BACTERIAL -GENE MUTATIONS) - ADDITION

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

Purity: 98-99 %



METHOD

Guideline followed: OECD 409, Edition 1 (Based on those described by Ames et al., (1975) and Green and Muriel (1976). Also based on OECD 471/472, updated and adopted July 21, 1997. Type: *Salmonella-Escherichia coli*/ Mammalian-Microsome Reverse Mutation Assay

System of testing: Bacterial and Mammalian

GLP: Yes.

Year study performed: 2000

Species/strain: Escherichia coli: WP₂uvrA

Salmonella typhimurium TA98, TA100, TA1535, TA1537, TA 1538

Metabolic activation: Strains were treated in the presence and absence of metabolic activation by S-9 mix. (S9 homogenate was purchased from Molecular Toxicology, Inc, Batch 1022. The homogenate was prepared from male Sprague-Dawley rats that had been injected (ip) with Arochlor 1254 (200 mg/ml in corn oil) at 500 mg/kg, as described by Ames et al 1975.

Range – Finding (µg/plate): 6.67, 10, 33.3, 66.7, 100, 333, 667, 1000, 3330, and 5000.

Concentrations tested ($\mu g/plate$): 100, 333, 1000, 3330, and 5000.

Positive controls:

TA98: 2.5µg benzo(a)pyrene w/ S9, 1.0µg 2-nitrofluorene w/out S9

TA100: 2.5µg 2-aminoanthracene w/ S9, 2.0µg sodium azide w/out S9

TA1535: 2.5µg 2-aminoanthracene w/ S9, 2.0µg sodium azide w/out S9

TA1537: 2.5µg 2-aminoanthracene w/ S9, 2.0µg ICR-191 w/out S9

WP2uvrA: 25µg 2-aminoanthracene w/ S9, 1.0µg 4-nitroquinoline-N-oxide w/out S9

Cytotoxicity: A minimum of three non-toxic doses were required to evaluate assay data.

Isonox 132 was evaluated for the ability to induce reverse mutations either in the presence or absence of mammalian microsomal enzymes at the histidine locus in the genome of several strains of *Salmonella typhimurium* and at the tryptophan locus of *Escherichia coli* starin WP2uvrA

The results of the initial mutagenicity assay were confirmed in an independent experiment.

RESULTS

Under the conditions of this study, Isonox-132 did not cause a positive increase in the mean number of revertants per plate with any of the tester strains either in the presence or absence of microsomal enzymes prepared from Aroclor TM induced rat liver (S9).

CONCLUSIONS

The test material is not genetically active in the Salmonella and Escherichia coli assay.

DATA QUALITY

Study conducted according to OECD guidelines.

Study GLP Compliant.

REFERENCES

Covance Laboratories Final Report on "Salmonella – Escherichia Coli/ Mammalian – Microsome Reverse Mutation Assay with a confirmatory Assay with Isonox-132." Covance Study No. 21109-0-409OECD. Unpublished.

15.2 GENETIC TOXICITY IN VITRO (CHROMOSOMAL ABERRATIONS) - ADDITION

TEST SUBSTANCE

4-sec-Butyl-2,6-di-tert-butylphenol

CAS No. 17540-75-9

Purity: 98-99%

METHOD

OECD Guideline No 437, Edition No. 3.

Type: Chromosome aberration assay System of testing: Non bacterial

GLP: Yes.

Year study performed: 2000

Species/strain: Chinese hamster ovary (CHO) cells

Metabolic activation: S-9 mix (with and without)

Concentrations tested:

In the initial trial of this assay, replicate culture of CHO cells were incubated with: 4.56, 6.52, 9.31, 13.3, 19.0, 27.2, 38.8, 55.4, 79.1, 113, 162, 232, 331, 473, and 675 μ g/ml for 3 hours with and without S9 and harvested at 20.0 hours after the initiation of treatment.

Cultures treated with 6.52, 9.31, 13.3, and 19.0 μ g/ml without metabolic activation and 4.56, 6.52, 9.31, and 13.3 μ g/ml with metabolic activation were analysed for chromosomal aberrations.

In the confirmatory trial, replicate cultures of CHO cells were incubated with 0.157, 0.313, 0.625, 1.25, 2.50, 5.00, 7.50, 10.0, 12.5, 15.0, and 20.0 μ g/ml without S9 and 2.50, 5.00, 7.50, 10.0, 12.5, 15.0, and 20.0 μ g/ml with S9. Treatment periods of 17.8 and 3 hours were used with and without S9, respectively, and cultures harvested at 20 hours after the initiation of treatment.

Cultures treated with 7.50, 10.0, 12.5, and 15 μ g/ml without S9 and 10.0, 12.5, 15.0, and 20.0 μ g/ml with S9 were analysed for chromosomal aberrations.

Controls: In the non-activation assays, negative controls were cultures containing only cells and culture medium. Vehicle controls were cultures containing DMSO, the vehicle for the test article, at

 $10 \mu g/ml$, the highest concentration used in test cultures. In the activation assays, the negative and vehicle controls were the same as described in the non-activation assays but with the S9 activation mix included.

The positive control agents which were used in the assays were mitomycin C (MMC) for the non-activation series and cyclophosphamide (CP) in the metabolic activation series. MMC is a clastogen and does not require metabolic activation. CP does not act directly but must be converted to active intermediates by microsomal enzymes. In the chromosomal aberrations assay, two concentrations of MMC (0.750 and 1.50 μ g/ml, initial assay, and 0.200 and 0.400 μ g/ml, confirmatory assay) and CP 95.00 and 10.0 μ g/ml) were used to induce chromosomal aberrations in the CHO cells. One of the concentrations was analysed in each of the aberration assays. Both MMC and CP were dissolved in water.

The *in vitro* metabolic activation system consisted of a rat liver post-mitochondrial fraction (S9) and an energy-producing system (NADP plus isocitric acid). Various hepatic P450 isoenzyme levels are increased by treatment of the rats with Aroclor TM 1254 (single concentration of 500 mg/kg) and sacrificed 5 days later. The S9 fraction, prepared in sucrose or KCl, was retained frozen at = 70 C until use.

The test article was dissolved in DMSO and treated at 1% (10.0 μ l/ml). A stock solution of Isonox 132 was prepared as a concentration of 67.5 mg/ml for the initial assay. The vehicle control cultures were treated with 10 μ l/ml of DMSO. The high dose tested in the initial assay, 675 μ l/ml, was above the solubility limit of Isonox 132 in the culture medium.

RESULTS

No increase in cells with chromosomal aberrations was observed at the concentrations analysed in the initial or confirmatory trials.

CONCLUSIONS

Isonox-132 was considered negative for inducing chromosomal aberrations in CHO cells with and without metabolic activation.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

Covance Laboratories Report on "Chromosomal Aberrations Test of Isonox-132 in Chinese Hamster Ovary (CHO) Cells." Covance Study No. 21109-0-4370ECD. Unpublished.

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): No.

Year study performed: 1949.

RESULTS

Melting point: 131°C.

CONCLUSIONS

The test substance has a melting point of 131°C.

DATA QUALITY

Not a GLP study.

Classed as handbook data by the Beilstein database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Sears, K., J. Amer. Chem. Soc., 71 (1949) as quoted in the Beilstein database.

2.1 BOILING POINT

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1944.

RESULTS

Boiling point: 278°C.

Pressure: atmospheric.

CONCLUSIONS

The test substance has a boiling point of 278° C.

DATA QUALITY

Not a GLP study.

Classed as handbook data by the Beilstein database.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Pardee, W., Ind. Eng. Chem., 36 (1944) as quoted in the Beilstein database.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: information not available in database.

GLP (Y/N): no.

Year study performed: 1984.

RESULTS

Vapor pressure: 0.000661 mm Hg @ 25°C (0.088 Pa).

CONCLUSIONS

The test substance has a vapor pressure of : 0.000661 mm Hg @ 25°C.

DATA QUALITY

Not a GLP study.

Purity of the test substance / decomposition: information not available in database.

REFERENCES

Perry, R.H. and Green, D. (1984); Perry's Chemical Engineers' Handbook., 6^h ed. McGraw-Hill, New York as quoted in SRC PhysProp database.

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4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: not known

GLP (Y/N): not known.

Year study performed: 1992.

RESULTS

Log Kow: 6.06.

CONCLUSIONS

The test substance has a log Kow of 6.06.

DATA QUALITY

Temperature and test substance purity not available.

Information taken from SRC PhysProp Database.

REFERENCES

Chem Inspection and Testing Institute; Biodegradation and Bioaccumulation Data of Existing Chemicals; Japan Chemical Industry Ecology-Toxicology and Information Centre. ISBN 4-89074-101-1 (1992) as quoted in the SRC PhysProp Database.

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: calculation using a log Kow value of 6.39.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: $0.267 \text{ mg/l} @ 25^{\circ}\text{C}$.

CONCLUSIONS

The test substance has a calculated water solubility of 0.267 mg/l @ 25°C.

DATA QUALITY

Calculation method

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

The log Kow value (6.39) calculated using the programme KOWWIN (see 4.1) has been used for the calculation of water solubility.

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $14.4 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 8.0 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 8.0 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

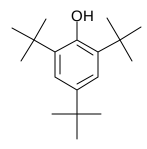
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3



COMMENT

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:262Data temperature (°C):25Water solubility (mg/l):0.267Vapor pressure (Pa):0.088Log Kow:6.06Melting point (C):131

Environmental conditions: defaults used.

RESULTS

Environmental compartment	percentage of test substance
Air	1.65
Soil	96.1
Water	0.095
suspended sediment	0.067
fish	0.0054
Sediment	2.13

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in days - weeks

Ultimate biodegradation in weeks - months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade slowly.

DATA QUALITY

Calculation method.

REFERENCES

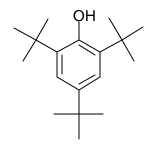
SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Fish

Exposure period: 96 hours, 14 days, 30 days, 90 days

RESULTS

LC50 (96hr) 0.076 mg/l

ChV (30 day) 0.011 m g/l

ChV (90 day) 0.003 mg/l

Remark: log Kow used 6.39 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.076 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

Calculation performed for this HPV submission

11.1 ACUTE TOXICITY TO ALGAE

TEST SUBSTANCE

2,4,6-Tri-tert-butylphenol

CAS No. 732-26-3

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.017 mg/l

ChV (96hr) 0.021mg/l

Remark: log Kow used 6.39 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.017 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

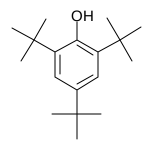
Calculation performed for this HPV submission

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2001

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.226 mg/l

ChV (21day) 0.008 mg/l

Remark: log Kow used 6.39 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.226 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

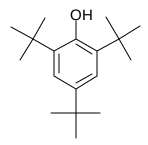
Calculation performed for this HPV submission

HEALTH ELEMENTS

13.1 ACUTE TOXICITY ORAL

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol



CAS No. 732-26-3

Purity of the test substance: Specific value not given but claimed to be of high purity.

METHOD

Method: Not specified, but based on EU and US EPA guidelines.

GLP: Data not available

Year study performed: 1989

Species/strain: Wistar rat

Sex: assumed 5 males and 5 females

No. of animals per sex per dose: 10

Vehicle: The test substance suspended in olive oil

Route of administration: Oral

Dose level: Not specified. 10 animals were used for each dosing and at least 6 increasing dose levels were selected and tested.

RESULTS

The LD50s were calculated using the Litchfield and Wilcoxon method.

LD50 male: 1670 mg/kg/body weight (95% confidence limits 1507 – 1851)

LD50 female: 1610 mg/kg/body weight (95% confidence limits 1423 – 1821)

Clinical signs included sedation and ataxia. Necropsy findings revealed swelling of liver and adrenal.

CONCLUSIONS

The liver was identified as a target organ.

DATA QUALITY

Method claimed to follow EC and US EPA protocol.

REFERENCES

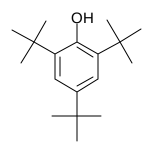
Acute Toxicity to 113 Environmental Chemicals.

Science Reports of the Research Institutes - Tohoku University, Series C: Medicine Vol. 36, Nos. 1-4, 1989.

16.1 REPEATED DOSE TOXICITY 24-month dietary

TEST SUBSTANCE

2,4,6-Tri-tert-butylphenol



CAS No. 732-26-3

Purity of the test substance: 97%.

METHOD

Method: Not specified

Test type: 24 months

GLP: Data not available

Year study performed: 1991

Species/strain: Slc: Wistar rats

Route of administration: Oral, feed

Exposure period: 24 months

Dose/concentration: 0, 30, 100, 300 and 1000 ppm. The highest dose was equivalent to approximately

1/20 of the LD50 value (1670 mg/kg in male) obtained from a preliminary acute toxicity test.

Sex: Males and females

No of animals per sex per dose: 40 males and 40 females

Frequency of treatment: Continuous in the diet.

Control group and treatment: Yes, concurrent, no treatment

Post exposure observation period: None

Statistical methods: One-way analysis of variance (ANOVA) with Dunnett's multiple comparison procedure.

Test conditions: Four-week old Wistar rats were divided into the 5 groups. The animals were fed diet containing the test substance for 24 months. General condition and body weights were observed and recorded throughout the experiment. Hematological and histopathological examinations were performed for all groups.

RESULTS

NOAEL 30 ppm (slight changes of hemoglobin concentration (Hb), mean corpuscular volume (MCV), blood urea nitrogen (BUN), glutamate oxaloacetate transaminase (GOT), phospholipids (PL), total cholesterol (T-Chol) and γ -glutamyl transpeptidase (γ -GTP) levels, but with liver and kidney weights within normal ranges, were found at this level).

Clinical signs:

Statistically significant reduction in body weight gain was found in female animals of the 1000 ppm group, 12 months after the start of the experiment and thereafter. No remarkable general findings and changes in food consumption were observed in the control and treated groups throughout the experimental period. Mortality in treated rats was comparable to that of controls.

Hematological and serum biochemistry findings:

6 months: Statistically significant decreases of Hb and MCV and significant increases of platelet count (Plt) (except in animals receiving 300 ppm), PL and T-Chol values were observed in male groups receiving 300 and 1000 ppm. Statistically significant decreases of MCV and GOT and statistically significant increases of T-Chol and γ -GTP values were observed in all treated female groups. Statistically significant decreases of Hb in the 300 and 1000 ppm groups and BUN in the 30 and 1000 ppm groups, and statistically significant increases of Plt in the 300 and 1000 ppm groups and γ -GTP in the 1000 ppm group were also found.

12 months: In males, statistically significant decreases of Hb in the 1000 ppm group, MCV in the 30, 300 and 1000 ppm groups and γ -GTP in the 30 and 100 ppm groups were observed. Levels of BUN and GOT in the 300 and 1000 ppm groups and PL and T-Chol in the 1000 ppm groups were significantly elevated. In females, statistically significant decreases of Hb in the two higher groups, MCV in the three higher groups, GOT in all treated groups and γ -GTP in the 100 ppm group were observed. Statistically significant increases of Plt and PL in the 100, 300 and 1000 ppm groups and T-Chol in all treated groups were found.

18 months: In males, statistically significant decrease of MCV in the 1000 ppm group, and significant increases of BUN in the 100 and 300 ppm groups and γ -GTP in the 1000 ppm group were observed. In all treated female groups, statistically significant decreases of Hb (except the 100 ppm group) and MCV, and statistically significant increases of PL and T-Chol were observed. Statistically significant decreases of GOT in the 30 and 300 ppm groups and γ -GTP in the 30 ppm group, and statistically significant increases of Plt in the 300 and 1000 ppm group were also found.

24 months: In males, statistically significant decreases of MCV in the 1000 ppm group, and significant increases of Plt in the 1000 ppm group and PL and T-Chol in the 100, 300 and 1000 ppm groups were observed. In females, statistically significant decreases of MCV in the 300 and 1000 ppm groups and γ -GTP in the 1000 ppm group, and statistically significant increases of Plt in the 1000 ppm group and PL and T-Chol in all treated groups were found.

Organ weight changes:

Statistically significant increases of relative liver weight in male rats receiving 300 and 1000 ppm and in all female treated groups except the 30 ppm at 12 months were found throughout the study. Statistically significant increases of kidney weights were observed in the animals receiving 1000 ppm at 12 months, in females receiving 1000 ppm at 18 months and in the males receiving 1000 ppm and females receiving 100, 300 and 1000 ppm at 24 months. Statistically significant increases of adrenal weights in the higher dose groups were also observed.

Histopathological findings:

In the 300 and 1000 ppm groups, swelling, focal necrosis and vacuolisation of liver cells were noted from 6 months after the start. No other statistically significant histopathological changes including neoplastic lesions were observed in other organs throughout the study.

Remarks:

In a previously carried out 3-month toxicity study, 15 male and 15 female Wistar rats were fed 0, 586, 1760 and 5280 ppm of TTBP (Matsumoto $et\ al.$, unpublished data). One week after the start of this experiment, marked reduction of body weight gain was noted in animals receiving 1760 and 5280 ppm, with some of the rats rapidly becoming moribund. In these cases, hematological, biochemical and histopathological examinations performed on days 8 to 14 revealed macrocytic anemia, focal necrosis of liver cells and hemorrhage in almost all organs. An increased value of T-Chol, transaminases and γ -GTP were noted in animals receiving 1760 and 5280 ppm, further indicating hepatotoxicity induced by TTBP.

Macrocytic anemia, reduced RCB, PCV and Hb values and inversely elevated MCV were observed following a massive dose of TTBP (Matsumoto and Shirai, 1987).

In another study, administration of 5600 ppm of TTBP in the diet for 7 days to male rats caused a significant reduction in blood coagulability, measured as prothrombin time, activated thromboplastin time and plasma recalcification time (Matsumoto *et al.*, unpublished data). These overall findings would suggest that the changes observed were due to the secondary effects following liver injury.

CONCLUSION

Administration of TTBP caused focal necrosis, swelling and vacuolisation of liver cells, increases in liver weight, slight microcytic anemia and elevation of serum phospholipid and cholesterol levels, presumably occurring as secondary effects following hepatotoxicity. The changes observed in female rats were more severe than those seen in males. Incidences of tumours in TTBP treated groups were not significantly higher than that of control.

DATA QUALITY

Reliable, without restrictions.

Although the test method and GLP data are not available, the test procedure closely follows the current testing guidelines.

REFERENCES

Matsumoto, K., Ochiai, T., Sekita, K., Uchida, O., Furuya, T. and Kurukawa, Y. (1991): Chronic Toxicity of 2, 4,6-Tri-*tert*-butylphenol in Rats. The Journal of Toxicological Sciences, **16**, 167-179 Matsumoto, K. and Shirai, T. (1987): Basic Study on Drug-induced Myelotoxicity: An Application of Bone Marrow Testing to Toxicity Study. Juntendo Med. J., **33**, 74 – 86 (in Japanese)

16.2 REPEATED DOSE TOXICITY 11 days toxicity in beagle dogs

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

Purity of the test substance: Data not available

METHOD

Method: Not specified

Test type: 11 days

GLP: Data not available

Year study performed: 1981

Species/strain: Beagle dogs

Route of administration: oral, feed.

Exposure period: 11 days

Dose/concentration: 0, 49.2, 173, 454 mg/kg/day.

Sex: Males

No of animals per sex per dose: 2 males per group.

Frequency of treatment: Daily

Control group and treatment: Yes, concurrent, no treatment

Post exposure observation period: None

Test conditions: Ten months old beagle males were given a daily dose of 2,4,6-tri-tert-butylphenol mixed with the dog food.

RESULTS

The food consumption rate was reduced in the high dose group. Diarrhoea was observed in two animals receiving 173 mg/kg and one receiving 454 mg/kg for 3 to 5 days. Faeces of one dog receiving

173 mg/kg and one receiving 454 mg/kg contained blood. Two animals of the high dose group showed signs such as gait abnormality (struggling), blepharoptosia, and decreased body temperature. The animals continued showing these signs until the end of dosing. From this observation, it was concluded that TTBP at this dosage has some effect on the central nervous system (apparently autonomic, principally).

Hematological examination was conducted on Day 11.

Parameters included glutamic -oxalacetic transaminase (GOT), glutamic -pyruvic transaminase (GPT), alkaline phosphatase (ALP), cholinesterase (CHE), total protein (TP), glucose (Glc), total cholesterol (CHO), urea-nitrogen (UN), sodium (Na), potassium (K), calcium (Ca), white blood cells (WBC), red blood cells (RBC), hemoglobin (Hgb) and hematcrit (Hct).

GOT, GPT and ALP increased relative to dose levels (marginal at 49.2 mg/kg).

CONCLUSION

TTBP did not cause hemorrhage or blood coagulation disorder to dogs. Behavioural abnormalities and increase in GOT, GPT and ALP were observed in animals of the highest dose group. Hematological observations indicated that TTBP might affect the liver parenchymal cells and increase metabolising enzymes.

DATA QUALITY

Methods/guideline – data not given

Purity of the test substance- data not given

REFERENCES

Toxicity of 2,4,6-tri-*tert*-butylphenol in Beagle Dogs, Tokyo Metrop. Res. Lab. Public Health, Tokyo. 32-2, 67-68, 1987.

OTHER TUMOUR INHIBITORY EFFECTS OF PHENOLIC COMPOUNDS ON BENZO(A)PYRENE-INDUCED NEOPLASIA

TEST SUBSTANCE

2,4,6-Tri-*tert*-butylphenol

CAS No. 732-26-3

Purity of the test substance: 97%

METHOD

Method: See below

GLP: Data not available

Year study performed: 1980

Species/strain: Mouse ICR/Ha strain

Route of administration: Oral, dietary

Exposure period: 38 days

Dose/concentration: 0.03 mmol/g (= 4,500 ppm in diet)

Sex: Females

No of animals per dose: 15

Frequency of treatment: continuous in the diet for 5.5 weeks. 1 mg benzo(a)pyrene (BP) by gavage, twice/week for 4 weeks. from second week of test diet administration.`

Positive control: BP as before, but no dietary addition of the test substance.

Post exposure observation period: 16 weeks

Statistical methods: Student's T test was used to determine the statistical differences in the number of tumors per group between the control and treated groups, and the χ^2 test was used for the differences in percentage of tumour bearing animals in these groups.

Test conditions: The mice were placed on a diet containing the test substance or diet without additions (control group) when they were 9 weeks of age. On the eighth day, the mice were given the first of 8

doses (2 times a week for 4 weeks) of 1 mg of BP in 0.2 ml of corn oil. The experimental diets were fed during the entire period of carcinogen administration and were discontinued 3 days after the last dose of carcinogen, at which time the mice were 98 days old. The mice were then fed pellets of rat chow, until they were killed at 211 days old and autopsied. Tumors of the forestomach were counted under a dissecting microscope. Tumors that were 1 mm or larger were recorded and checked histologically.

RESULTS

In the control group (animals given BP but no dietary addition of phenol), 90 - 100% of animals had neoplasms of the forestomach and the average number of tumours per animal was between 4.1 and 5.8. In the animals given dietary addition of 2,4,6-tri-*tert*-butylphenol, 93% of animals had neoplasms of the forestomach and there was no reduction in the number of neoplasms per animal.

CONCLUSION

Substitution of a third *tert*-butyl group *para* to the hydroxyl group apparently destroyed any inhibitory activity of 2,4,6-tri-*tert*-butylphenol in this mouse model, as compared with the di-substituted phenols assessed in this study, which did show inhibitory activity.

DATA QUALITY

Reliable, without restrictions

REFERENCES

Department of Laboratory Medicine and Pathology, University of Minnesota, Minneapolis. Cancer Research 40, 2820- 2823. August 1980.

PHYSICAL/CHEMICAL ELEMENTS

1.1 MELTING POINT

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: information not available.

GLP (Y/N): no.

Year study performed: not stated.

RESULTS

Melting point: 65℃.

CONCLUSIONS

The test substance has a melting point of 65°C.

DATA QUALITY

Not a GLP study.

Decomposition: information not available.

Information taken from a technical datasheet.

Purity given as 95% minimum.

REFERENCES

Schenectady International Inc. Technical datasheet.

1.2 MELTING POINT

TEST SUBSTANCE

2,4- Bis(alpha, alpha- dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: information not available.

GLP (Y/N): no.

Year study performed: 1950.

RESULTS

Melting point: 65.5 - 67°C.

CONCLUSIONS

The test substance has a melting point range of 65.5 - 67°C.

DATA QUALITY

Not GLP.

Information taken from a literature search covering appropriate databases.

Information on purity, decomposition and method not available.

REFERENCES

Patent: Goodrich Co., US 2714120 1950.

2.1 BOILING POINT

TEST SUBSTANCE

2,4- Bis(alpha, alpha- dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: Schenectady International Inc. internal procedure.

GLP (Y/N): No.

Year study performed: not stated.

RESULTS

Boiling point: >300°C at 760 mm Hg.

CONCLUSIONS

The test substance has a boiling point of >300°C at 760 mm Hg.

DATA QUALITY

Not GLP.

Information taken from a technical datasheet.

Decomposition: information not available.

Purity given as 95% minimum.

REFERENCES

Schenectady International Inc. Technical datasheet.

3.1 VAPOR PRESSURE

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Vapor pressure: $5.84 \times 10^9 \text{ mm Hg } @ 25^{\circ}\text{C} (7.8 \times 10^7 \text{ Pa}).$

CONCLUSIONS

The test substance has a calculated vapor pressure of $5.84 \times 10^{-9} \text{mm Hg} \ @ 25^{\circ}\text{C}$.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. MPBPWIN v 1.30

4.1 PARTITION COEFFICIENT

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: calculation.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Log Kow: 6.73.

CONCLUSIONS

The test substance has a log Kow of 6.73.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. KOWWIN v 1.63.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

5.1 WATER SOLUBILITY

TEST SUBSTANCE

2,4- Bis(alpha, alpha- dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: calculation method using a calculated log Kow of 6.73.

GLP (Y/N): no.

Year study performed: parameter value calculated for this HPV submission.

RESULTS

Water solubility: 0.0548 mg/l at 25°C.

CONCLUSIONS

The test substance has a calculated water solubility of 0.0548 mg/l at $25 \,^{\circ}\text{C}$.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. WSKOW v 1.33.

OTHER

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 (Mackay, D., Varhannickova, D., Ma, Kuo-Ching & Shiu, Wan-Ying, Chemosphere, Vol 29, No. 6, pp 1155-1224, 1994).

The log Kow value (6.73) calculated using the programme KOWWIN (see 4.1) has been used for this calculation of water solubility.

ENVIRONMENTAL FATE AND PATHWAY ELEMENTS

6.1 PHOTODEGRADATION

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: calculation using the programme AOPWIN v1.88.

Test type: calculation of the rate constant for the atmospheric reaction between photochemically produced hydroxyl radicals and the test substance in the vapor phase.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS

Sensitizer: hydroxyl radical.

Overall hydroxyl rate constant: $57.4 \times 10^{-12} \, \text{cm}^3/\text{molecule-sec}$.

Half-life: 2.24 hours.

CONCLUSIONS

The programme estimates that in a typical atmosphere 50% of the test substance will undergo reaction in 2.24 hours.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. AOPWIN v 1.88.

OTHER

No experimental data was found on direct aqueous photolysis of the test substance. However, *p*-cresol, a related substance, in aqueous solution is reported as having a half-life of 35 days in sunlight (Smith, J.H. et al, "Environmental Pathways of Selected Chemicals in Freshwater Systems: Part II. Laboratory Studies," EPA-600/7-78-074, May 1978. Cited in Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw-Hill, Inc., Washington, 1990, page 8-38.

7.1 STABILITY IN WATER

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

COMMENT

No abiotic hydrolysis studies were located.

The category phenols do not possess any functional groups that are regarded as being susceptible to hydrolysis under environmental conditions (Lyman, W.J., Reehl, W.F. and Rosenblatt, D.H., Handbook of Chemical Property Calculation Methods, McGraw -Hill, Inc., Washington, 1990, pages 7-4 and 7-5).

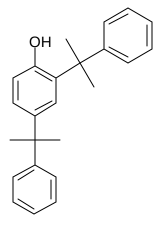
The software prediction programme HYDROWIN v1.66 cannot estimate hydrolysis rate constants for phenols.

8.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS (FUGACITY)

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4



METHOD

Test type: Calculation of partitioning between environmental compartments.

Year study performed: Model run for this HPV submission.

Model: Level 1 Fugacity-Based Environmental Partitioning Model v2.11.

Input values

Chemical specific

Molecular mass:330Data temperature (°C):25Water solubility (mg/l):0.0548Vapor pressure (Pa): 7.8×10^{-7} Log Kow:6.73Melting point (°C):65

Environmental conditions: defaults used.

RESULTS

Environmental compartment	Percentage of test substance
Air	1.95×10^{-5}
Soil	97.7
Water	0.021
suspended sediment	0.068
fish	0.0055
Sediment	2.17

DATA QUALITY

The Mackay Level I Fugacity Model estimates the equilibrium distribution of a fixed quantity of a non-reacting chemical in a closed environment at equilibrium; with no degradation reactions and no flow or intermedia transport processes. The chemical is assumed to distribute instantaneously to an equilibrium concentration and therefore the medium receiving the emission is unimportant. This model is an aid to understanding the physical chemistry properties that are of greatest importance in determining the environmental distribution of substances; it is not a tool to predict actual or likely concentrations in a real environment.

Alkylphenols are weak acids with typical pKa values in the range 9.9 to 10.9 and therefore their physical chemistry properties are unlikely to be affected by the pH values normally found in the environment.

REFERENCES

This software program is available with the publication: Mackay, D., Multimedia environmental models: the fugacity approach, Lewis Publishers Inc., Chelsea, MI, 1991.

9.1 BIODEGRADATION

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Method/guideline followed: calculation using the programme BIOWIN v3.65.

Test type: calculation of the probability for rapid aerobic biodegradation of the test substance in the presence of mixed populations of environmental microorganisms.

GLP (Y/N): no.

Year study performed: calculation made for this HPV submission.

RESULTS & CONCLUSIONS

The program predicts:

Primary biodegradation in weeks

Ultimate biodegradation in months.

This material is considered to be inherently biodegradable, and would aerobically biodegrade very slowly.

DATA QUALITY

Calculation method.

REFERENCES

SYRACUSE Chemical Properties Prediction Program. BIOWIN v 3.65.

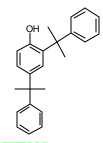
ECOTOXICITY ELEMENTS

10.1 ACUTE TOXICITY TO FISH

TEST SUBSTANCE

2, 4Bis (alpha, alpha-dimethylbenzyl) phenol

CAS No. 2772-45-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Fish

Exposure period: 96 hours, 30 days and 90 days

RESULTS

LC50 (96hr) 0.059 mg/l

ChV (30 day) 0.008mg/l

ChV (90 day) 0.003 mg/l Remarks: log Kow used 6.73 (calculated value)

CONCLUSIONS

Estimated LC50 (96hr) for the test substance was found to be 0.059 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

OTHER

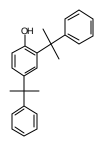
Calculation performed for this HPV submission.

11.1 TOXICITY TO AQUATIC PLANTS (E.G., ALGAE)

TEST SUBSTANCE

2, 4-Bis (alpha, alpha-dimethylbenzyl) phenol

CAS No. 2772-45-4



METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999. Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Green algae

Exposure period: 96 hours

RESULTS

EC50 (96hr) 0.011 mg/l

ChV (96hr) 0.016 mg/l

Remark: log Kow used 6.73 (calculated value)

CONCLUSIONS

Estimated EC50 (96hr) for the test substance was found to be 0.011 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v.0.99e

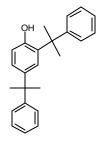
OTHER

Calculation performed for this HPV submission.

12.1 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G., DAPHNIA)

TEST SUBSTANCE

2, 4Bis (alpha, alpha dimethylbenzyl) phenol



CAS No. 2772-45-4

METHOD

ECOWIN Classes for Microsoft Windows. US EPA, March 1999.

Estimated method ECOSAR v 0.99e

GLP: (N)

Year study performed: 2000

Species: Daphnid

Exposure period: 48 hours, 21 days

RESULTS

LC50 (48hr) 0.21 mg/l

ChV (21day) 0.007 mg/l

Remark: log Kow used 6.73 (calculated value)

CONCLUSIONS

Estimated LC50 (48hr) for the test substance was found to be 0.21 mg/l.

DATA QUALITY

Estimation

REFERENCES

ECOSAR v. 0.99e

OTHER

Calculation performed for this HPV submission.

HEALTH ELEMENTS

13.1 ACUTE TOXICITY SKIN IRRITATION

TEST SUBSTANCE

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

METHOD

Acute Dermal Irritation OECD Guidelines No. 404 (1981)

GLP: (Y)

Year study performed: 1991

Species: New Zealand White rabbits

Number of animals: 3

A single, 4-hour, semi-occluded application of the test material was applied to the intact skin of three rabbits.

Vehicle: 0.5 g of the test material was ground and moistened with 0.5 ml of distilled water; introduced under a 2.5 cm x 2.5 cm gauze patch and placed in position on shorn skin. The patch was secured by surgical adhesive tape. Corsets were used to prevent the animal from interfering with the patches. Four hours after application, the corset and patches were removed, and residual material was carefully removed.

The test sites were examined at 1 hour following the removal of patched and 24, 48 and 72 hours later.

RESULTS

Very slight erythema was noted at all treated skin sites at 1 hour, and at two treated skin sites at the 24-hour observation. Very slight edema was noted at 2 treated skin sites at 1 hour. All treated skin sites appeared normal after 24-48 hours.

CONCLUSIONS

The test material produced a primary irritation index of 0.3 and was classified as a mild irritant according to the Draize classification scheme. No corrosive effects were noted.

DATA QUALITY

Study conducted to GLP and OECD guidelines.

REFERENCES

2,4-DCP-Acute Dermal Irritation Test in the Rabbit, Safepharm Laboratories Ltd, P.O. Box No. 45, Derby, DE1 2BT, U.K.

OTHER

Study Ref: 47/1555

HEALTH ELEMENTS

2,4- Bis(alpha, alpha-dimethylbenzyl)phenol

CAS No. 2772-45-4

No additional test reports found